



מדינת ישראל
STATE OF ISRAEL

REC'D 22 JUL 2004

WIPO

PCT

Ministry of Justice
Patent Office

משרד המשפטים
לשכת הפטנטים

This is to certify that
annexed hereto is a true
copy of the documents as
originally deposited with
the patent application
particulars of which are
specified on the first page
of the annex.

זאת לתעודה כי
רצופים בזה העתקים
נכונים של המסמכים
שהופקדו לכתחילה
עם הבקשה לפטנט
לפי הפרטים הרשומים
בעמוד הראשון של
הנספח.

PRIORITY DOCUMENT
SUBMITTED OR TRANSMITTED IN
COMPLIANCE WITH
RULE 17.1(a) OR (b)

This

05-07-2004

היום

רשם הפטנטים

רשם הפטנטים

Commissioner of Patents



נתאשר
Certified

160987	מספר: :Number
21-03-2004	תאריך: :Date
חוקדם / נדחה: :Ante / Post-dated	

בקשה לפטנט

Application for Patent

אני, (שם חמבקש, מענו - ולגבי גוף מאוגד - מקום התאגדות)

I (Name and address of applicant, and, in case of body corporate place of incorporation)

Disc-O-Tech Medical Technologies, Ltd.
3 Hasadnaot Street
Herzelia 46728
Israel

דיסקוטק טכנולוגיות רפואיות בע"מ
רח' חסדנאות 3
הרצליה 46728
ישראל

ששמה הוא Law
Of an invention, the title of which is

בעל אמצאה מכח חדין
Owner, by virtue of

כלים ושתלים הניתנים לעיוות

(בעברית)
(Hebrew)

DEFORMABLE TOOLS AND IMPLANTS

(באנגלית)
(English)

Hereby apply for a patent to be granted to me in respect thereof

מבקש בזאת כי ינתן לי עליה פטנט

*בקשת חלוקה - Application of Division		*בקשת פטנט מוסף - Application for Patent Addition		*דרישה דין קדימה Priority Claim	
מבקשת פטנט from Application No. _____ מס' _____ Dated _____ מיום _____		*לבקשה/לפטנט to Patent/Appl. No. _____ מס' _____ Dated _____ מיום _____		מספר / סימן Number/Mark	תאריך Date
ייפוי כח: כלל/מיוחד - רצוף בזה / עוד יוגש P.O.A: general / individual - attached / to be filed later חוגש בעניין _____ filed in case _____					
המען למסירת הודעות ומסמכים בישראל Address for Service in Israel פנסטר ושות' קניין רוחני 2002 בע"מ רח' בזל 16 פ"ת ת.ד. 10256 פ"ת, 49002					
עבדור המבקש, חתימת המבקש Signature of Applicant פנסטר ושות' קניין רוחני 2002 בע"מ		שנת 2004 Of the year		בחודש מרץ Of	היום 21 This
				לשימוש הלשכה For Office Use	

110/03694

טופס זה, כשהוא מוטבע בחותם לשכת הפטנטים ומושלם מספר ובתאריך ההגשה, הינו אישור לחגשת הבקשה שפרטיה רשומים לעיל.
This form, impressed with the Seal of the Patent Office and indicating the number and date of filing, certifies the filing of the application, the particulars of which are set out above.

Delete whatever is inapplicable ימחק את המיותר

כלים ושתלים הניתנים לעיוות

DEFORMABLE TOOLS AND IMPLANTS

דיסקוטק טכנולוגיות רפואיות בע"מ

Disc-O-Tech Medical Technologies, Ltd.

c:110/03694

DEFORMABLE TOOLS AND IMPLANTS**RELATED APPLICATIONS**

The present application claims the benefit under 35 USC 119(e) of 60/478,841 filed on June 17, 2003 and US Provisional Application for Patent named "Expandable Disc Nucleus Prosthesis filed December 2003, the disclosures of which are incorporated herein by reference.

The present application is related to US 09/890,172 and US 09/890,318, the disclosures of which are incorporated herein by reference.

FIELD OF THE INVENTION

The present invention is related to tools and implants which are deformed during use, for example for orthopedic use.

BACKGROUND

Spinal compression fractures are painful and disfiguring fractures in which a spinal vertebra axially compresses. This causes distortion and shortening of the spine and also pain, like other fractures. A conventional treatment is bed rest, which heals the fracture but does not reset the bone in its original configuration.

US patent application publication 2002/0156482A1 Scribner et al., the disclosure of which is incorporated herein by reference, describes a procedure by which a balloon is inserted into a spinal vertebra and then expanded, in an attempt to reset the vertebra to its original configuration. Thereafter, bone filler may be injected into a void created by the balloon, to fixate the bone.

PCT publication WO 00/44319, the disclosure of which is incorporated herein by reference, suggests a device which can be mechanically expanded to support two vertebra or to be inserted inside a bone to support a fracture.

SUMMARY OF THE INVENTION

An aspect of some embodiments of the invention relates to a pliable medical element (hereafter a "deformer") configured to deform from filling a narrow diameter volume to filling or defining a larger diameter volume. In an exemplary embodiment of the invention, the deformer comprises a slotted tube, formed of a polymer or fabric material, such that when the tube is axially compressed, a plurality of leaves extend to the sides of the tube and fill the volume surrounding the tube. Optionally, a locking element, for example an inner bolt or wire is used to maintain the tube in its deformed configuration.

In an exemplary embodiment of the invention, the material used is pliable enough to provide a defomer with some degree of conforming to a body geometry, for example cortical

plates of a vertebra, while still being able to apply a force against an external resisting body (such as the plates). This is in contrast to cement, which simply flows to where there is least resistance, even though cracks and out of the vertebra. This is also in contrast to a balloon, which, being filled with a fluid, tends to equalizes pressures on different parts thereof, thus
5 force is applied in a direction of least resistance.

In an exemplary embodiment of the invention, a deformer is used to apply a desired degree of deformation displacement, while a balloon can apply a force, but not determine a displacement. One potential disadvantage of some implementations of a balloon is that expanding a balloon near a vertebral wall may cause application of force on the wall and
10 fracturing thereof. A deformer, in accordance with exemplary embodiments of the invention can be deformed with a small safety margin, for example, 1 mm, 2, mm, 3mm, or less, from the vertebra.

In an exemplary embodiment of the invention, a deformer is used for a spinal application, for example as a tool for expanding a compressed vertebra and/or as an implant for
15 spacing between or inside a vertebra.

In an exemplary embodiment of the invention, a deformer is deformed using the following process. A distal end of the deformer is (optionally) held in place while a proximal end of the deformer is pushed or pulled axially towards the distal end. An overtube surrounding the deformer is optionally provided to control the deformation process. Optionally, the overtube
20 is retracted in conjunction with the pushing (or pulling), to provide serial deformation of the deformer. One potential advantage of the optional holding of the distal end of the deformer, is that the deformer can thus deform in place without retracting from tissue.

The deformer can have various shapes once deformed, for example, the deformer can be cylindrical, cone line (with a truncated tip) or lordotic (a truncated 4 sided pyramid). The
25 deformer may or may not have rotational symmetry around the axis of the tube. For example, the deformer may be concave on one side. In another example, the defomer bends the axis when deforming. In another example, the cross-section of the deformer is non-uniform and/or rotationally asymmetric.

Optionally, the pliability and/or other mechanical properties, such as flexural modulus, tensile strength and/or elongation of the deformer vary along its length and/or at different
30 angles thereof.

In an exemplary embodiment of the invention, neighboring leaves of the deformer support each other when the deformer is deformed. Optionally, the leaves at one or both ends of

the deformer are shorter, so that they can make do with support from only one side of leaves. Optionally alternatively or additionally, the end leaves are made less pliable. Optionally alternatively or additionally, the end leaves are designed to extend axially and not only radially as in some other designs.

5 Optionally, the leaves of a deformer substantially fill the space outside the deformer, in the volume defined thereby, for example, 30%, 40%, 50%, 60%, 70%, 80%, 90% or more of the volume excluding a volume of an inner rod.

 While a slotted tube is described, in other embodiments, other base forms are used for the deformer. For example, an unslotted tube may be used. In another example, slots are not
10 through the entire thickness of the tube.

 In an exemplary embodiment of the invention, the deformer is used as an implant. Optionally, the implant is delivered on a delivery tube and then released. In an exemplary embodiment of the invention, the deformer implant includes two end plates that are interconnected by a bar or wire. The end plates, optionally made of metal, are optionally used
15 to apply compressive force in an axial direction against the ends of the pliable ends of the deformer. This type of implant may be used, for example, in a spinal vertebra, or for supporting fallen end plates in long bones.

 Optionally, the implant includes one or more radio-opaque markers and/or the end plates act as such markers.

20 In an exemplary embodiment of the invention, the deformer is used as a void creating and/or tissue moving tool. In an exemplary embodiment of the invention, the deformer is mounted on a rod, one end of which rod is fixed to a first end of the deformer and another end of which rod is coupled to an element that pushes a second end of the deformer towards the first end. One or more radio-opaque markers may be provided.

25 In an exemplary embodiment of the invention, the deformer is used as a tissue engaging element. In one example, the deformer is used to hold an implant inside a medullar channel of a bone, for example, for holding a prosthesis, for holding an intra-medullar nail or for holding a femoral head screw. In another example, two deformers are used, each one holding a different tissue section and optionally interconnected by a bar, wire or hinge.

30 In an exemplary embodiment of the invention, while being deformed, the volume taken up by the deformer is reduced, for example, due to compression of the deformer material (e.g., made of a porous, fabric or compressible material), or due to compression of voids formed in the material. Optionally alternatively or additionally, the deformer is compressed into an inner

channel thereof (e.g., a lumen of a tube deformer), so that the total external volume defined by the deformer is reduced, for example, by 5%, 10%, 20% or more.

Optionally, the deformer has a composite structure. In one embodiment of the invention, one or more threads, for example of metal or Kevlar are embedded in the implant, for example, to increase tensile strength and/or to modify mechanical properties of the deformer. Optionally alternatively or additionally, the deformer is composed of axial segments that are welded or otherwise attached, each segment having different properties.

Optionally, the deformer is configured to elute a material with biochemical properties, for example, a bone growth enhancing material. Optionally, the deformer is coated with such a material or a delivery vehicle for the material. Alternatively or additionally, for example, the deformer is impregnated with the material and/or includes one or more void filled with such material, so that when deformed, the material is eluted. Optionally, the eluted material is a cement or a cement hardener.

An aspect of some embodiments of the invention relates to a method of treating a spinal compression fracture or a fallen bone plate in a long bone, using a deformer. In an exemplary embodiment of the invention, a deformer is guided into a vertebra, for example using a guide wire and then deformed so that the vertebra is axially expanded. Possibly, an old fracture can be re-broken using this method. Then, the deformer is removed and cement or bone slurry is injected and used to set the vertebra. Optionally, a balloon or leaky balloon or fabric bag is inserted to hold at least some of the cement and the cement is injected into the balloon. Optionally, the balloon is biodegradable, to allow bone growth through it. In an alternative embodiment, the deformer is left in place as an implant. Optionally, cement is injected through the deformer. Optionally, a smaller amount of cement is required and there is optionally a reduced danger of leakage. Optionally, the implant is biodegradable, at least in part. In an alternative embodiment of the invention, the deformer is provided inside a balloon or fabric bag and used to expand the vertebra. Cement or bone slurry are optionally used to replace or as an addition to the deformer, which is optionally removed.

In an exemplary embodiment of the invention, a pliable deformer is used. Alternatively or additionally, a stiff implant, for example made of titanium, is used.

In an alternative vertebral treatment method, a deformer is implanted into an inter-vertebral space, to serve as a cushion or disc replacement between two vertebrae.

An aspect of some embodiments of the invention relates to a system for deforming a deformer. In an exemplary embodiment of the invention, the system includes a rod on which

the deformer is mounted and attached at a distal end thereof. A pusher tube pushes a second end of the deformer towards the distal end. An overtube retracts so that selected parts of the deformer are unrestrained to expand. Optionally, a first part of the implant is exposed and the pullback of the overtube is delayed until that first part deforms. Optionally, the pushing is mechanically coupled to retraction of the overtube and includes a mechanical delaying mechanism.

In an exemplary embodiment of the invention, a last delivery step of the deformer comprises tightening the deformer so that its radial resistance increases. In an exemplary embodiment of the invention, this is achieved by axially compressing the deformer a final amount. Possibly, this allows a final radial force to be applied in concert by an elongate section of the deformer, rather than by short sections one at a time.

In a non-implant configuration, the system optionally includes a mechanism for controlling the length of deformed deformer. In one example, retraction of the overtube is stopped when a desired length is achieved.

In an exemplary embodiment of the invention, the deformer is cannulated to allow cement flow therethrough. Alternatively or additionally, the system itself and/or the rod on which the deformer is mounted, cannulated to allow such flow. Optionally, the rod includes a lumen, for example, for a guide wire.

In an implant configuration, the system optionally includes a mechanism for locking and releasing a deformer implant.

Optionally, the system can be used for un-deforming the deformer, for example, by pulling back on the pusher, which is optionally attached to the proximal end of the deformer. Optionally, such un-deforming does not return the overtube to its original position, thereby preventing reuse of the system.

Optionally, the system is flexible, for example for use in an endoscope. Optionally, a pull wire is used rather than a pusher, to deform the deformer. For example, the pull wire can be mounted on a pulley to pull the proximal part of the deformer towards its distal part.

Optionally, a hydraulic mechanism is used for pushing the pusher and retracting the over tube. Optionally, a hydraulic column is used to push the proximal end of the deformer and deform it.

An aspect of some embodiments of the invention relates to a balloon system for pushing apart end plates of a vertebra. In an exemplary embodiment of the invention, the system comprises an outer balloon and an inner balloon. The inner balloon is inflated first, pushing

apart the plates and/or also limiting the expansion of the outer balloon to at least include the shape of the inner balloon. It is expected that the inner balloon should be able to, in most cases expand properly, thereby preventing migration of the outer balloon and/or setting a limit on the sideways extent of the outer balloon. Optionally, the balloons are inflated in a staggered manner. Optionally, a third or additional balloons are provided enclosing the first and second balloons.

An aspect of some embodiments of the invention relates to applying a desired displacement inside the body. Optionally, the displacement is applied with some degree of compliance to body geometry, while not allowing the displacement to be diverted to a different direction as a balloon might. Optionally, the displacement is applied with an increase in diameter of a displacing device, for example, of at least 50%, 100%, 150%, 200%, 300%, or any intermediate or greater value.

In an exemplary embodiment of the invention, the displacement is applied in one direction while a sensitive tissue lies in a second direction. Optionally, the displacement is directed to not damage the sensitive tissue.

BRIEF DESCRIPTION OF THE FIGURES

Particular embodiments of the invention will be described with reference to the following description of exemplary embodiments in conjunction with the figures, wherein identical structures, elements or parts which appear in more than one figure are optionally labeled with a same or similar number in all the figures in which they appear, in which:

Fig. 1A is a schematic illustration of an undeformed deformer, in accordance with an exemplary embodiment of the invention;

Figs. 1B-1H are schematic illustrations of a deformer during deformation, in accordance with an exemplary embodiment of the invention;

Figs. 2A-2E are schematic axial cross-sectional views showing the deformation of the deformer of Fig. 1A, in accordance with an exemplary embodiment of the invention;

Figs. 3A-3E show steps in the treatment of a vertebra, in accordance with an exemplary embodiment of the invention;

Fig. 4A is a flowchart of a method of treating a vertebra in accordance with Figs. 3A-3E;

Figs. 4B-4H are parts of a kit for treating a vertebra, in accordance with an exemplary embodiment of the invention;

Figs. 5A-5D are schematic views of a delivery system, in accordance with an exemplary embodiment of the invention;

Fig. 6 is a schematic cross-sectional view of a hydraulic delivery system in accordance with an exemplary embodiment of the invention

5 Fig. 7A is a schematic cross-sectional view of a belt based delivery system in accordance with an exemplary embodiment of the invention

Figs. 7B and 7C illustrate a wire based delivery system, in accordance with an exemplary embodiment of the invention;

10 Figs. 7D and 7E illustrate a soft-material based delivery system, in accordance with an exemplary embodiment of the invention;

Figs. 8A-8C illustrates an implant release and locking mechanism, in accordance with an exemplary embodiment of the invention;

Figs. 9A-9F illustrate various deformer geometries in accordance with exemplary embodiments of the invention;

15 Fig. 10 schematically shows a spinal joint in accordance with an exemplary embodiment of the invention;

Figs. 11 and 12 shows the use of an implanted deformer for supporting a humerus head (Fig. 11) and a tibial plateau (Fig. 12), in accordance with an exemplary embodiment of the invention.

20 Fig. 13 is a schematic illustration of an intra-medullar nail, in accordance with an exemplary embodiment of the invention;

Fig. 14 illustrates a hip trochanter support implant using a deforming element in accordance with an exemplary embodiment of the invention;

25 Figs. 15A and 15B illustrates a dental implant in accordance with an exemplary embodiment of the invention; and

Figs. 16A-16E show a balloon-in-balloon configuration, in accordance with an exemplary embodiment of the invention.

DETAILED DESCRIPTION OF EXEMPLARY EMBODIMENTS

Slotted Tube Expander Design

30 Fig. 1A is a schematic illustration of an undeformed deformer 100, in accordance with an exemplary embodiment of the invention. Fig. 1F is a schematic illustration of a fully deformed deformer 100, in accordance with an exemplary embodiment of the invention.

In this design, deformer 100 comprises a tube body 102, having a plurality of slots 104 formed along its length, in an axial direction. In the embodiment shown, slots 104 are arranged in alternating lines 106 and 108 such that a plurality of alternating leaf lines are formed, comprising leaves 110 alternating (axially and radially) with leaves 112. As will be described below, for example, other designs may be provided. A distal end 114 and a proximal end 116 are also marked on Fig. 1A.

As can be seen in Fig. 1F, when fully deformed, deformer 100 substantially fills a volume of space. Also, as shown, the leaves support each other.

Figs. 2A-2E are schematic axial cross-sectional views showing the deformation of the deformer of Fig. 1A, in accordance with an exemplary embodiment of the invention.

Fig. 2A shows deformer 100 mounted on a delivery rod 202 and optionally attached (e.g., at distal end 114 thereof) to an end 204 thereof. A pusher 206, for example a tube, pushes proximal end 116 of deformer 100 towards distal end 114. An overtube 208 restrains the radial deformation of deformer 100, as will be described below.

In Fig. 2B, pusher tube 206 is advanced, while overtube 208 is not retracted. As a result, a first plurality of leaves 210 extend radially past overtube 208.

In Fig. 2C, pusher tube 206 is advanced more (for example continuously or in discrete steps) while overtube 208 is retracted (for example continuously or in (optionally matching) discrete steps). The ratio between the movements can be axially linear or non-linear, for example. As a result, a second plurality of leaves 212 extend radially past overtube 208 and optionally lean on leaves 210. As can be seen, leaves 210 and 212 are not axially compressed to the fullest extent possible. In some embodiments, as more leaves are extended out, the tips of previously extended leaves move axially. Optionally, this axial motion is used for engaging tissue or engaging a nearby deformer, for example for dual deformer use.

Fig. 2D shows the state after all the leaves have been deformed out. It should be noted that the process of leaf deformation may appear in some embodiments like an extrusion, even though the deformer is not flowing.

Fig. 2E shows the effect of further axial pushing by pusher 206 once all the leaves have been extruded. The result shown is an axial compression and radial stiffening of the leaves. Optionally, this simultaneous radial stiffening of multiple locations along the deformer, allows force to be applied to nearby tissue in a desired direction.

Referring back to Fig. 1F, it should be noted that in some cases not all of the leaves are extended in a perfect manner. However, in many applications this is not a problem. In some

applications, the pliability of the material allows the other leaves to adjust their position accordingly. In some applications, the imperfect extension allows space for cement and/or other fluids.

Referring in more detail to Figs. 1B-1H. Fig. 1B shows deformer 100 before any leaves
 5 are extended. Fig. 1C shows deformer 100 after four sections of leaves are extended. Fig. 1D shows deformer 100 after some more leaves are extended. Fig. 1E shows deformer 100 with nearly all the leaves extended. Fig. 1F shows deformer 100 with all the leaves extended and with axial compression.

Fig. 1G shows deformer 100 during removal, due to retraction of pusher 206. Fig. 1H
 10 shows deformer 1H fully un-deformed so it can be removed from the body.

In some embodiments of the invention, deformer 100 is deformed all at once, without an overtube. Suitable treatment or design of the deformer may, however, impose an order on the deformation.

Vertebral Treatment system

15 In an exemplary embodiment of the invention, a deformer is used to treat compression fractures of the vertebrae, by inserting a deformer in a narrow diameter configuration, into a vertebra and deforming the deformer such that it expands the vertebra.

Figs. 3A-3E show steps in the treatment of a vertebra, in accordance with an exemplary embodiment of the invention. Fig. 4A is a flowchart 400 of a method of treating a vertebra in
 20 accordance with Figs. 3A-3E. Figs. 4B-4H are parts of a kit for treating a vertebra, in accordance with an exemplary embodiment of the invention. Other kits may include fewer or greater number of elements.

Referring to Fig. 4A. At 402, a vertebra 300 (Fig. 3A) is accessed using a Jamshidi needle 430 (Fig. 4B), with an inner stylet or guidewire.

25 At 404, a guide wire 432 (Fig. 3B) is guided through needle 430 into vertebra 300.

At 406, needle 430 is removed (Fig. 3B). As can be appreciated, other methods of accessing a vertebra can be used and many are known in the art. In addition, it is not essential to use a guidewire for accessing the vertebra and other means can be used to guide deformer 100 to its target.

30 At 408, a cannula 440 (Fig. 4C), with an optional trocar 442 (Fig. 4D) are guided along guidewire 432 to vertebra 300. A handle 444 (Fig. 4C-1) optionally couples the cannula and the trocar and/or can be attached to the drill or other tools (e.g., a modular handle). Cannula 440 is optionally rotated to ensure it engages vertebra 300 and/or nearby bones. Other fixation

methods, for example, forward pressure or sideways extending leaves or a balloon may be used to hold cannula 440 in place.

At 410, guide wire 432 and/or trocar 442 are removed, leaving only cannula 440.

At 412, a drill 450 (Fig. 4E) is used to ream out a section of vertebra 300 and form a void 302 (shown in Fig. 3C) for deformer 100. Optionally, the drill is guided along guidewire 432.

At 414, a biopsy is optionally taken. Optionally, a biopsy is taken earlier, for example at 402 or 404.

At 416, a dummy tool 460 (Fig. 4F) is optionally inserted and x-rays are acquired to check its placement (Fig. 3C). Other verification means may be used as well. Optionally, the verification is used to select a deformer diameter and/or length.

At 418, deformer 100, mounted on a delivery system 500 (Fig. 5) is inserted into void 302 (Fig. 3D). It should be appreciated that a deformer can be inserted into a vertebra from various directions. Further, multiple deformers can be inserted into a same vertebra from multiple direction, for example posteral or lateral approaches and/or from either side of the body plane, in either approach.

At 420, deformer 100 is deformed (Fig. 3E). During such deformation, a determination of the actual deformation of vertebra 300 may be acquired, to ensure correct expansion and/or prevent over expansion and/or fracturing damage to vertebra 300. For example, x-ray or CT images may be used.

At 422, deformer 100 is optionally removed and a cement delivery tube 470 (Fig. 4G) is inserted instead. Delivery tube 470, in one embodiment, includes an outer tube section 472 and an inner plunger 474. Cement, stored in a lumen of outer tube 472 is forced by the plunger into void 302. Exemplary materials which may be injected include, bone chips, bone slurry (e.g., auto-graft, xenograft, allograft, from cadavers), PMMA, calcium phosphate and/or calcium sulfate.

At 424, all the tools are removed and the procedure is completed. Surgical holes, etc. maybe closed using methods known in the art.

In an alternative procedure, deformer 100 is released at the end of act 420 and remains in the body. Optionally, deformer 100 is mounted on a cannulated rod, through which cement or other materials can be provided to void 302. Alternatively or additionally, the cement is provided via an over tube (not shown) which surrounds rod 202 (Fig. 2A) and has a diameter small enough to reach into the vertebra, so cement will not leak out.

In an alternative procedure, cement delivery tube 470 is replaced with a balloon delivery element 480 (Fig. 4H), which expands a balloon 482 inside void 302, to maintain the shape of vertebra 300. The expansion can be, for example, with cement, with a fluid, such as saline and/or with particle matter, such as bone fragments. Balloon 482 is optionally biodegradable in the body, for example being made of Poly(L-Lactide-co-capralactone) 70:30 or Poly(L-Lactide-co-glycolide) 85:15, 82:18 or 10:9. Optionally, a mesh is used instead of a balloon, to allow leakage of some bone cement.

In an alternative procedure, a balloon and deformer delivery system is used, in which a deformer deforms inside a balloon. Optionally, cement and/or other materials are provided through a channel provided in the delivery system.

Details of Exemplary Deformer Delivery System

Figs. 5A-5D illustrate a deformer delivery system 500 in accordance with an exemplary embodiment of the invention. Fig. 5A shows delivery system 500 before deformation of deformer 100. Fig. 5B shows system 500 after deformation of deformer 100.

System 500 comprises a body 502 having a handle 504 for deforming deformer 100. In other embodiments, a power source, for example, an electric motor or a hydraulic power source are used instead of handle 504. When handle 504 is rotated, a rod 505 attached thereto is rotated. A threading 506 on rod 505 engages a matching threading or projection on a nut 508. Nut 508 is coupled to pusher 206, for example, via a pin 510. Thus, rotation of handle 504 advances pusher 206. It should be noted that in an exemplary embodiment of the invention, rod 202 does not move and maintains its end 204 in a fixed relationship to body 502

The rotation of rod 505 optionally retracts overtube 208, as well. Optionally, the retraction is delayed relative to the advancing of pusher 206. In an exemplary embodiment of the invention, the following mechanism is used. A nut 524 is mounted on a distal part of rod 505, with a threading 522 which optionally has a smaller pitch than threading 506, so retraction of overtube 208 is less pronounced than advancing of pusher 206. Overtube 208 is coupled to a block 512 that is moved by nut 524. However, in an exemplary embodiment of the invention, once retracted, overtube 208 cannot return to its starting position. For example, a ratchet mechanism may prevent such motion. Optionally, a locking disk 515 allows only one way motion of block 512, relative to pusher 206.

Meeting of nut 508 and block 512 optionally stops the deforming of deformer 100. Alternatively, one or more stops may be provided to prevent motion of one or both of nuts 508 and 524 and/or block 512. Optionally, such stops are movable, for example to define various

deformer lengths. Optionally, the threading on the nuts is flexible enough (or frangible) to allow rod 505 to rotate while the nut is held in place.

Optionally, a pin 528 in nut 524 extends outside of body 502 and serves as a marker on a scale, to show a status and/or degree of deformation.

5 Optionally, one or more markings 526 are provided on overtube 528, for example, to indicate its retraction degree and/or to assist in determining a depth in the body.

 In an exemplary embodiment of the invention, inadvertent motion of overtube 208 during insertion into the body is prevented using a locking mechanism. Figs. 5C and 5D shows a detailed view of an exemplary locking mechanism in operation, in which motion of a nut
10 allows a lock holding the overtube in place to be released and allows the block to be moved, optionally by the same nut.

 In Fig. 5C, a lock 514 locks block 512 to a nub 516 in body 502. A bottom surface 530 of nut 524 prevents lock 514 from moving out of the way, by pressing against an upper surface 532, thereof. This prevents inadvertent motion of block 512 (and overtube 208), for example
15 during insertion.

 In Fig. 5D, once nut 524 retracted sufficiently, lock 514 is not blocked and will disengage from nub 516 when block 512 is retracted by nut 524, for example by an inclined surface 538 of lock 514 sliding past an inclined surface 536 of nub 516.

 It should be noted that a same delivery system can be used for devices where deformer
20 100 stays in the body and devices where deformer 100 is part of system 500.

 Optionally, tensioning state is provided in which after deformer 100 is deployed, additional motion of pusher 206 is provided without motion of overtube 208, for example, to tighten deformer 100. In one example, threading 522 ends in a manner that allows free rotation relative to nut 524, but no axial advance (e.g., a stop), without a corresponding end to threading
25 506. In another example, all of rod 506 is moved axially, which also may include motion of block 512, if suitably coupled to rod 506.

 Optionally, threadings 522 and/or 506 are non-uniform, for example, to provide a certain non-linear relationship between the motions of pusher 206 and overtube 208.

 Optionally, system 500 is used to inject cement or another material into the vertebra. In
30 some embodiments of the invention, cement is injected after deformer 100 is removed or after system 500 is removed. Alternatively, cement is provided through system 500. In an exemplary embodiment of the invention, rod 202 is hollow and cement is provided from a cement source

513 through a tube 511 connected to rod 202. Optionally, rod 202 is aperture underlying deformer 100 and/or at its distal end 204.

Alternative Delivery Systems

While linear compression of deformer 100 is shown, in other embodiments, rotational
5 motion is provided, for example, by pusher 204 rotating relative to rod 202, optionally being threaded on it.

Optionally, system 500 includes means to limit the forces applied to the vertebra. Optionally, a mechanical fuse is provided in system 500 so that if a certain force is exceeded, the fuse tears or slips and no further or greater force is applied. Optionally, pin 510 serves as
10 such a fuse. Optionally, a string (not shown) is used to retract nut 508 if pin 510 tears. Optionally, pin 510 is designed to bear 20 Kg without tearing. Greater or smaller forces, for example, 10 Kg, 50 Kg or 100 Kg, or smaller or greater forces may be provided as well.

Alternatively or additionally, a warning or indication display is used. In one example, a force sensor (not shown) is provided in rod 202 and which senses the force applied to it by
15 deformer 100. Alternatively or additionally, a strain sensor is provided on rod 202 to measure axial strain. Optionally, such sensors are wired to a warning LED or scale on body 502.

Fig. 6 shows a hydraulic powered delivery system 600, in accordance with an exemplary embodiment of the invention. Fluid (e.g., saline, oil or air) enters a chamber 604 via an inlet 602., pushing against a piston 606 which is free to move in a cylinder 608. Optionally,
20 excess fluid exits through an outlet port 610. Optionally, the hydraulic pressure is manually supplied, for example using a hand pump

Motion of piston 606 is coupled to pusher 204 via a coupler 612. Retraction of overtube 208 is optionally provided using a method as described in Fig. 7A, below.

Fig. 7A is a cross-sectional view of an alternative delivery system 700, in accordance
25 with an exemplary embodiment of the invention. A handle 702 is attached to a body 704 and can rotated relatively thereto. An optional power gear 706 reduces the motion of rotation, to increase its mechanical gain and turns a belt 708. A block 712 rides on the belt and pushes pusher 206. Optionally, belt 78 slips when it attempts to apply too great (and possibly dangerous) a force.

30 In an exemplary embodiment of the invention, the following mechanism is used to couple a retraction of overtube 208 with advance of pusher 204. A tongue 714 interconnects a pin 710 of block 712 with a pin 720 of a block 718 coupled to overtube 208. Prior to pin 710 reaching an inclined section 716 of tongue 714, a hook 722 prevents retraction of overtube 208.

Once inclined section 716 is reached, tongue 714 moves and pin 720 slides along an inclined surface 724 (shown as dashed). This sliding causes retraction of overtube 208.

Incline 716 and surface 724 can also be non-linear.

Figs. 7B and 7C illustrate a tension based delivery mechanism 780, in accordance with an exemplary embodiment of the invention. Instead of a pusher rod or tube 204, a disc 782 is provided which compresses deformer 100. In an exemplary embodiment of the invention, a wire 784 is attached to disc 782 at a point 788, travels along a lumen 786 along deformer 100 to distal end 214. Optionally, lumen 786 is formed between rod 202 and deformer 100. Optionally, lumen 786 is formed as a groove in rod 202.

At distal end 214, a curved lumen 790 turns wire 784 back towards disc 782. Optionally, wire 784 exits disc 782 via an aperture 794 therein.

In operation, when wire 784 is pulled back, disc 782 advances and deforms deformer 100.

In an alternative embodiment, disc 782 is advanced by pressure of a fluid, rather than by tension from a wire.

In an alternative embodiment, disc 782 engages rod 202 using a threading on rod 202 and/or disk 782. Advancing of disc 782 is optionally by rotation of disc 782.

It should be noted that the forces applied by deformer 100 may be relative small until the final compression. Thus, even a flexible delivery system is not expected to be deformed at least during most of the deformation process. Optionally, the delivery system is provided via an endoscope or is otherwise navigable. Optionally, such flexible delivery is used for delivering some of the implants described below. Optionally, overtube 208 is flexible and is inelastic enough to prevent undesired radial distortion of deformer 100 except where desired.

Optionally, one or more of rod 202, pusher 204 and overtube 208 are bent.

Figs. 7D and 7E illustrate an alternative deforming mechanism, in which a pliable material is distorted from a narrow diameter to a greater diameter.

In Fig. 7D, a portion 793 of soft material, such as silicon is contained within overtube 208. Optionally, portion 793 comprises a stiffer bag with an inner contents which are softer. A wire 796 optionally interconnects a distal tip 797 with a base section (not shown).

In Fig. 7E, a pusher tube 794 is advanced, forcing the silicon to have a radially expanded shape.

Optionally, one or more of the above delivery methods, systems and mechanism is used for delivering or deploying a cage device and/or other devices, for example as described in PCT

publication WO 00/44319 and WO 00/44321, the disclosures of which are incorporated herein by reference.

Variations on Vertebral Treatment

As noted above, in some embodiments of the invention, cement is injected while deformer 100 is inside the body or after it is removed. In some embodiments, the cement or another material is injected into a balloon/mesh, before or after deformer 100 is removed. In some embodiments, no cement or other material is injected.

In a particular variation, a deformer is removed and a metal cage device is inserted. This device may then be filled with materials such as cement or bone chips, as described herein.

A potential advantage of injecting cement after a deformer 100 is in place, is that as most of the volume is filled by the deformer, a smaller amount of cement is needed and there may be less danger of leakage or undesirable migration of the cement. Also, as a void is already created and held open, lower pressures may be used to advance the cement.

Another potential advantage is that the cement can be sweated out through deformer 100, thus possibly holding deformer 100 together and/or assist in providing a uniform distribution of cement. Optionally, pockets are formed for cement between different sets of leaves, which pockets are generally decoupled, so that increased pressure in one may not cause leakage from another one.

Figs. 8A-8C illustrate a deformer release mechanism 800 in accordance with an exemplary embodiment of the invention. Deformer 100 is mounted on a rod 802 that engages an extension 804 with a distal tip 806. In Fig. 8A, deformer 100 is undeformed. In Fig. 8B, a disc 812 has been advanced so that it deforms deformer 100 and engages a narrowing 808 in extension 804. For example, disc 812 can be super-elastic. Retraction of disc 812 is optionally prevented by a base 810 of extension 804.

In Fig. 8C, rod 802 is removed from extension 804, for example, by being unscrewed from a recess 814 which engages a tip thereof (not shown, for example using threading). To remove deformer 100, a disc 812 can be shape memory and a cooling fluid provided to make it pliable and easy to remove.

In an example of an automatic release mechanism, disc 812 when entering recess 812 shear one or more wires (not shown) which pass in recess 808 and attach extension 804 to rod 802.

Optionally, in some embodiments of the invention deformer 100 is a titanium or other metal device, for example as described in earlier applications of the assignee of the present invention.

Variations on deformer Design

5 A deformer may have various cross-sectional shapes, for example, being a rectangle, a square, a circle, an ellipse and/or a concave shape. Optionally, the cross-sectional shape and/or dimensions change along the axis of the deformer. Optionally, various shapes are achieved by suitable cutting of slot lengths. Alternatively or additionally, a deformer shape may be achieved by cutting the deformer after it is deformed to have the desired geometry and then un-
10 deforming the deformer. Optionally, the deformer is circuit based on 3D or 2D images of The cross-section may be, for example rotationally symmetric, mirror symmetric or asymmetric.

The axial shape may also be of various types, for example, uniform, lordotic (at one side a greater diameter than the other), ellipsoid (narrower diameter at both ends), hour-glass, or varying (e.g., diameter increases and decreases several times along its length).

15 The material from which the deformer is made can be, for example, of uniform thickness and uniform properties. Alternatively one or both of the thickness and material properties can vary, resulting, for example, in a deformer which has a non-uniform deformation. The varying can be, for example, in an axial and/or angular directions.

Optionally, properties of a deformer are made to vary by varying one or more of length,
20 direction, width, linearity, and/or spatial density of slots. Alternatively or additionally, alignments of pairs of slots (which define a leaf) is varied. For example, slots can have a helical pattern, be arranged in lines and/or vary in axial and/or radial densities.

Optionally, no slots are provided. For example, the deformer may be twisted or compressed (and is optionally made more elastic). Alternatively or additionally, the slots do not
25 reach through the thickness of the deformer, for example being only inside and/or only outside.

In an exemplary embodiment of the invention, one side of the deformer is made softer so that deformation will be preferable in a certain direction.

Fig. 9A shows six different exemplary axial profiles, hourglass (902), off-axis symmetric (904), ellipsoid (906), lordotic (908), inverse lordotic (910), and off-axis
30 asymmetric (912). Other off-axis designs can be used.

In a particular example, Fig. 9B shows a deformer 914, in which slots are formed on one side, resulting in deformation which is not symmetric with respect to the axis.

The contact area between the deformer and the tissue, can be, for example, bumpy (as shown in Fig. 1), or smooth, for example, if the device is encased in a bag, or if the ends of the leaves are treated so that they are softer than other parts of the deformer.

In many of the embodiments described, an inner rod is used to lock the deformer.
5 Optionally, a wire is used to lock the deformer by interconnecting the two ends of the deformer. Optionally, the ends fold in, at least slightly, so that they are protected from outside tissue, for example, so they do not contact bone.

Optionally, the inner bar is made of a super elastic or shape memory material that determines the final shape of the deformer, once released and/or during release. For example,
10 the inner rod can be pre-trained to achieve a curved or spiral shape, in 2D or in 3D.

Optionally, the deformer is shaped so that multiple deformer will interlock or fit side by side, for example, a concave depression in one deformer matching another deformer. Optionally, multiple deformers are implanted and/or deformed at a same time or even simultaneously.

15 Optionally, one or more leaves have defined thereon hairs or projections to engage tissue and/or encourage ingrowth or adhesion.

One property of deformers in accordance with some embodiments of the invention is that leaves are supported by leaves. As a result leaves at the ends lack some support. Optionally, the end leaves are made shorter. Alternatively or additionally, these end leaves are
20 made stiffer. Alternatively or additionally, these end leaves bend over the axis. Alternatively or additionally, a greater number of leaves and/or axial or radial leaf density are provided at the end.

Fig. 9C shows a proximal end 920 for attaching a deformer thereto, in accordance with an exemplary embodiment of the invention. End 920 comprises a tubular section 922 adapted
25 to engage a pusher 906, for example by contact, adhesive or threading. End 920 comprises an aperture section 924 including a plurality of apertures 924, on which proximal end 116 of deformer 100 can be mounted and melted on to ensure engagement.

Figs. 9D and 9E illustrate a distal end cap 931 for attaching a deformer thereto, in accordance with an exemplary embodiment of the invention.

30 An outer body 930 and an inner rosette 932 having a plurality of petals 934 define between them a lumen 938 into which distal end 114 of deformer 100 is inserted. Heat is then applied to melt the plastic into the metal. Alternatively or additionally, adhesive may be used. Optionally, an aperture 936 is provided for venting air, if required. Optionally, a lumen 940 is

defined through rosette 932 and body 930, and is optionally threaded to engage an end 214 of rod 202.

The length of a deformer (in a deformed state can vary depending on the application, for example, being between 2mm and 100 mm, for example, 10 mm, 20 mm, 30 mm, 40 mm, or any smaller, larger or intermediate values. A ratio of axial shortening, can be, for example, 1:2, 1:3, 1:4, 1:5, 1:8, 1:10, or any smaller, intermediate or greater ratio. A ratio of radial increase, can be, for example, 1:2, 1:3, 1:4, 1:5, or any smaller, intermediate or greater ratio. The radius of an undeformed deformer can be, for example, 1mm, 2mm, 3mm, 4mm, 5mm, 7mm, 10mm or any smaller, intermediate or greater radius. The thickness of the deformer material can be, for example, 0.5 mm, 1 mm, 2 mm, 3 mm, or any smaller, intermediate or greater thickness. Axial and/or radial leaf density per length unit can be, for example, 1:4 mm, 1:3mm, 1:2mm 1:1mm 1:0.5 mm or any smaller, intermediate or greater per unit density. Leaf length can be, for example, 1 mm, 3 mm, 5 mm, 7 mm, 10 mm, 20 mm, or any smaller, intermediate or greater length.

15 Materials

In an exemplary embodiment of the invention, the material used has a shore hardness between 50A and 90D, for example, 90A, 60D. Optionally, the material is selected so that individual leaves can be moved by the forces applied by the bone (e.g., a spine), while multiple leaves will be able to support each other against such forces.

20 In one example, a device as described above, with cylindrical diameter, deforms from a diameter of 5 mm to 15 mm, is 20 mm long, made from polyurethane, has a shore hardness 90A, is form of a tube 1.6 mm thick and was bench tested to lift 60 Kg.

In an exemplary embodiment of the invention, pliability and/or leaf density are selected to control a desired space filling effect.

25 The material used is optionally elastic.

Optionally, a compressible material is used. In one example, a fabric, for example Gore-Tex, Dacron or a metal mesh are used.

Optionally, a natural material, such as cotton or collagen are used.

30 Optionally, a tradeoff between material pliability and conformance of the device is achieved. For example, as the material is made softer, the device may be more conforming during implantation and possibly even after final tightening of the device.

Optionally, deformer 100 is coated with various materials, for example, an adhesive, osteo-conductive materials, growth promoters, anti-inflammatory, antibiotics, radioactive materials or other a materials known in the art..

Optionally, deformer 100 is made smooth to assist in its removal after a time.

5 Optionally, deformer 100 is made degradable so that it degrade after time and doe not need to be removed and/or is made partially degradable so that tissue in-growth can occur. Optionally, different parts degrade at different rates, for example, the inner locking bar degrading only after a long time or not at all.

10 In an exemplary embodiment of the invention, deformer 100 is formed of a composite material. In one example, deformer 100 is manufactured by stringing beads of various materials and then melting them together to form a tube. Alternatively, a deformer is made from segments which are merely strung together and possibly adhered to each other. Optionally, different beads have different mechanical and/or degrading properties.

15 Fig. 9F shows another type of composite device, 950, in which a plurality of wires, for example, Kevlar or metal wires 952 are embedded therein. Optionally, the wires are not embedded but are found in channels 954, optionally, allowing relative motion of the wires and deformer 950.

20 Alternatively or additionally, radially-directed wires may be embedded. Alternatively or additionally, a helical wire is provided. Optionally, one or more radial wires are provided at one or both ends of the deformer. Optionally, such end wires prevent tearing of deformer 950 and may optionally be locked together by a locking wire.

25 A potential advantage of using wires is to prevent tearing of the deformer. Another potential advantage of using wires is that a deformer may be removed by pulling on the wires, rather than on the deformer. Another potential advantage of using wires is changing material properties locally and/or providing strength where needed.

Artificial Disc Application

30 In an exemplary embodiment of the invention, deformer 100 is used as an implant for a disc. In an exemplary embodiment of the invention, some or all of the disc material is removed. Alternatively, no disc material is removed. A deformer 100 is then implanted inside the inter-vertebral space, to support and/or expand the existing disc. Optionally, an implant which curves or curls is used. Alternatively, two implants may be inserted side by side.

Artificial Joint

Fig. 10 shows vertebral joint 1000, in accordance with an exemplary embodiment of the invention. Joint 1000 comprises a first bone-engaging element 1002 engaging an inner volume of a vertebra 1004 and a second bone-engaging element 1006 engaging an inner volume of a vertebra 1007. The two elements 1002 and 1006 are interconnected by a bar or rod 1008, which either includes a hinge or serves as a living hinge between the vertebra. For example, bar 1008, can be flat so as to have a preferred bending direction. Alternatively or additionally, bar 1008 is notched on one side, so as to prefer bending in one direction rather than bending back.

In an exemplary embodiment of the invention, device joint 1000 is deployed in the following manner. A cannula is guided to vertebra 1004 and an aperture 1020 made therein. A dotted line 1026 shows a path of a flexible drill (not shown) that enters (and optionally forms) aperture 1020 and then forms an aperture 1022 in the base of vertebra 1004 and an aperture 1024 in a top of vertebra 1007. A guide tube is provided along this path. Joint 1000, in a narrowed diameter state is inserted along this path. A mechanism as described above is used to push a ring or slotted ring 1010 to axially compress engaging element 1006 against a base 1012, while retracting the guide tube. When the compression is completed, ring 1010 can lock. Then, the guide tube is further retracted and compression of element 1002, by a ring 1016 against a base 1014, proceeds. In an exemplary embodiment of the invention, advancing of ring 1010 (for compressing element 1006) is done relative to ring 1014. When completed, ring 1016 can lock. Optionally, a flexible delivery system is used. Alternatively, a hinge is provided at ring 1016.

In an alternatively embodiment of the invention, joint 1000 is used as a replacement finger joint. In this embodiment a substantially straight and rigid delivery system can be used.

Bone Implants

Figs. 11 and 12 shows the use of an implanted deformer 100 for supporting a humerus head (Fig. 11) and a tibial plateau (Fig. 12), in accordance with an exemplary embodiment of the invention.

In these embodiments, a deformer is inserted near the inside of an end plate of a long bone to support a sunken bone section. Optionally, the deformer is used on an old fracture, to reset the bone. Optionally, the deformer selected is positioned to press against two cortical bone sections, one stronger and one to be moved or supported. Alternatively or additionally, one side of the deformer may rest on compressed (but the deformer) spongy bone. Alternatively or additionally, the deformer is configured to expand radially more in one direction than in

another. Alternatively or additionally, the deformer is configured to be wider in one direction than another, so as to preferentially support motion of the deformer in the narrower direction.

Fig. 13 shows an intra-medullar nail 1300, in accordance with an exemplary embodiment of the invention. Nail 1300 includes deforming elements 1302 and 1304 at either end and a bar 1306 interconnecting them. Optionally, elements 1302 and 1304 degrade in the body so that only bar 1306 needs to be removed. Optionally, bar 1306 also degrades, but at a lower rate. A potential benefit of this design is that the medullar canal may remain mostly undamaged and/or free.

In an alternative nail design, elements 1302 and 1304 are formed of a continuous tube, slit only at the areas of the elements. Optionally, an over tube (not shown) sheaths this single tube between elements 1302 and 1304.

Fig. 14 shows a deformer 1400 in a femoral head application, for example for holding the bone together or for supporting a nail. Alternatively or additionally, a deformer may be used to hold a prosthesis, for example being placed in the femoral medullar channel to hold a hip implant. Optionally, one or more deformers are placed between a cortical bone and an implant, to apply compressive forces to one or both of the bone and the implant.

In a pedicle screw application, or in other applications, a deformer section may be used to apply force and anchor against spongy bone, in addition to or alternatively to hard cortical bone.

20 Dental Implant

Figs. 15A and 15B show a dental implant 1500 in accordance with an exemplary embodiment of the invention. One potential advantage of using a deformer-type implant is that in some embodiments an implant can be partially tightened, readjusted and then tightened some more. Optionally, once the implant is completely tightened, a cap is mounted on the implant.

Another potential advantage for dental usage is that a deformer may be able to better grip (and not apply too much force) against a weakened jaw bone (cortical and/or spongy bone). Possibly, the softer materials used in the implant prevent or slow down degradation by wear of the bone, as there is less of a point pressure applied.

Balloon Device

30 Figs. 16A-16E illustrate a balloon in balloon expansion device, in accordance with an exemplary embodiment of the invention.

Fig. 16A shows a balloon device 1600 inserted in a vertebra 300 with a fracture 1602. Device 1600 comprises an inner balloon 1604 and an outer balloon 1606.

Figs. 16B-16E are cross-sectional views along a line II-II, showing the operation of device 1600.

In Fig. 16B, device 1600 is not inflated.

5 In Fig. 16C, inner balloon 1604 is inflated, partially moving apart end plates 1608 and 1610 and/or fixing balloon 1606 in place.

In Fig. 16D, outer balloon 1606 is partially inflated. As can be seen, balloon 1606 is limited in its ability to reach the sides of vertebra 300, where it might cause damage.

In Fig. 16E, balloon 1606 is further inflated, further spacing apart vertebra end plates 1608 and 1610.

10 Optionally, inner balloon 1604 is replaced by a deformer, for example of the type described above.

It will be appreciated that the above described methods of implanting and treating may be varied in many ways, including, changing the order of steps, which steps are performed more often and which less often, the arrangement of elements, the type and magnitude of forces
15 applied and/or the particular shapes used. Further, the location of various elements may be switched, without exceeding the spirit of the disclosure, for example, the location of the power source. In addition, a multiplicity of various features, both of method and of devices have been described. It should be appreciated that different features may be combined in different ways. In particular, not all the features shown above in a particular embodiment are necessary in
20 every similar exemplary embodiment of the invention. Further, combinations of the above features are also considered to be within the scope of some exemplary embodiments of the invention. In addition, some of the features of the invention described herein may be adapted for use with prior art devices, in accordance with other exemplary embodiments of the invention. The particular geometric forms used to illustrate the invention should not be
25 considered limiting the invention in its broadest aspect to only those forms, for example, where a cylindrical tube electrode is shown, in other embodiments an rectangular tube maybe used. Although some limitations are described only as method or apparatus limitations, the scope of the invention also includes apparatus programmed and/or designed to carry out the methods.

Also within the scope of the invention are surgical kits which include sets of medical
30 devices suitable for implanting a device or material and such a device. Section headers are provided only to assist in navigating the application and should not be construed as necessarily limiting the contents described in a certain section, to that section. Measurements are provided to serve only as exemplary measurements for particular cases, the exact measurements applied

will vary depending on the application. When used in the following claims, the terms "comprises", "comprising", "includes", "including" or the like means "including but not limited to".

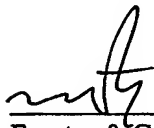
5 It will be appreciated by a person skilled in the art that the present invention is not limited by what has thus far been described. Rather, the scope of the present invention is limited only by the following claims.

CLAIMS

1. A deformer, comprising:
5 an axial member; and
a pliable tube mounted on said axial member and adapted to be deformed from a first narrower diameter to a second greater diameter.
2. A method of treating a bone, comprising:
10 inserting a unsealed pliable element into the bone; and
mechanically deforming the pliable element such that said pliable element applies deforming force on the bone.
3. A method according to claim 2, wherein said pliable element comprises at least one
15 open aperture of cross-section greater than 0.5x0.5 mm.

For the Applicant,

20



Fenster & Company, Intellectual
Property 2002 Ltd.
c:110/03694

25

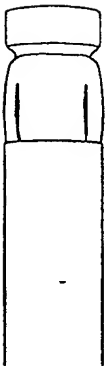


FIG. 1B



FIG. 1A

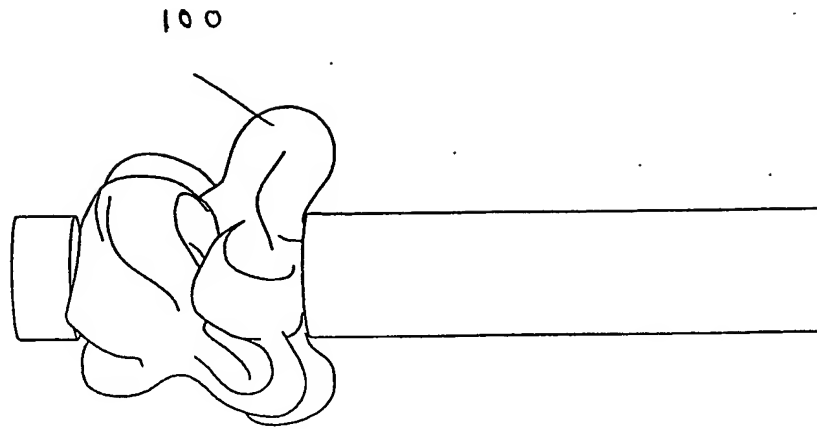


FIG. 1 C

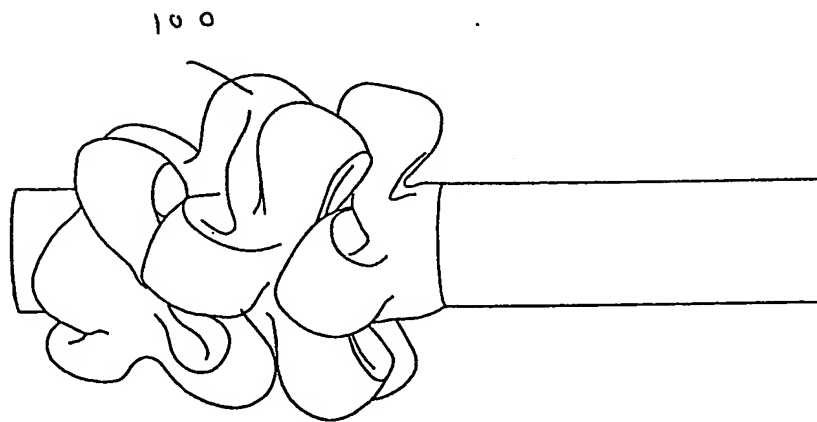


FIG. 1 D

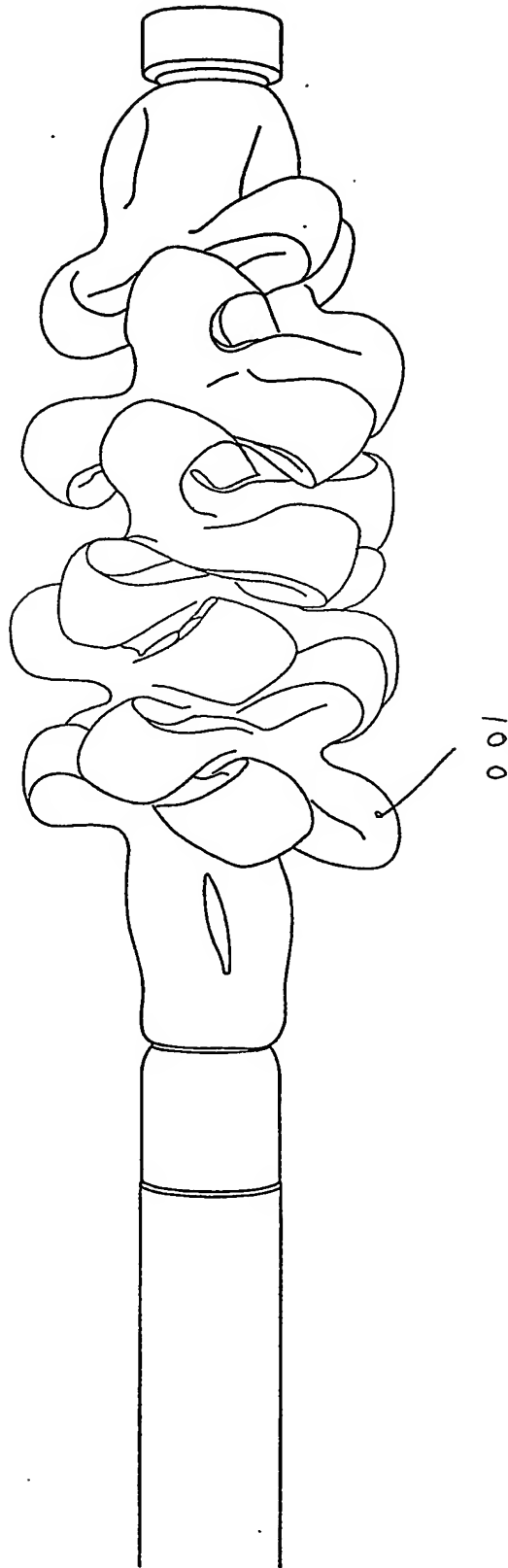


FIG. 1E

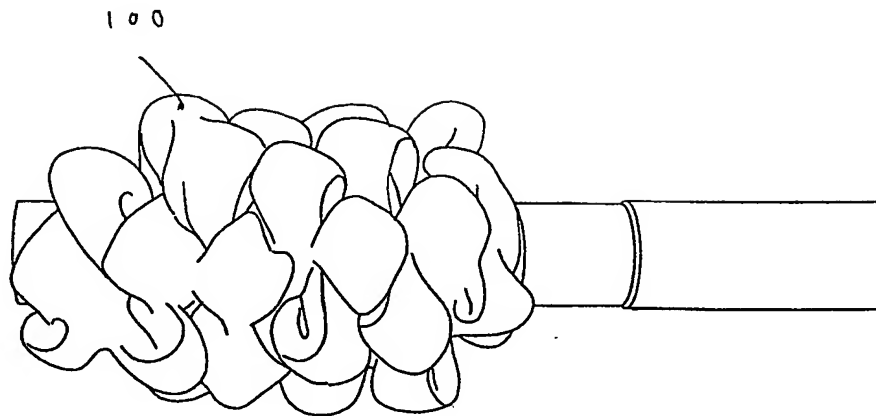


FIG. 1E

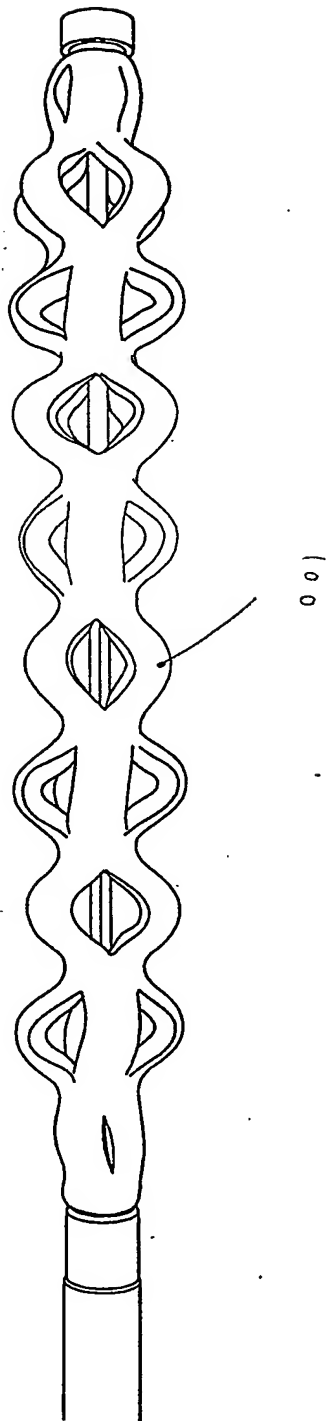


FIG. 16

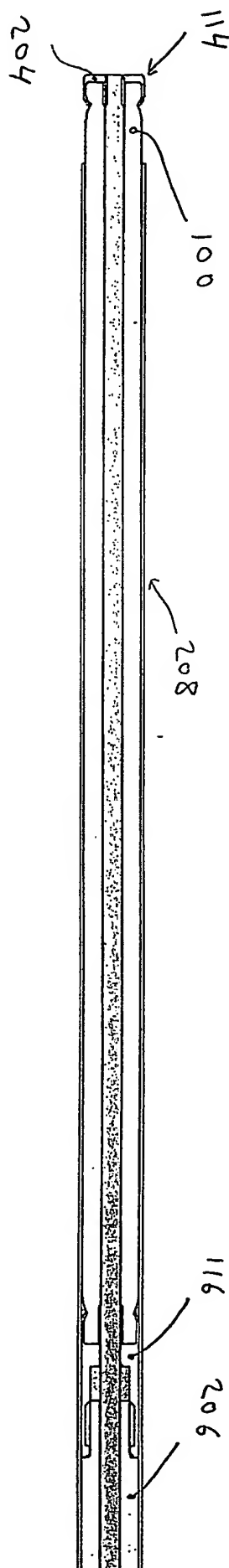


Fig. 2A

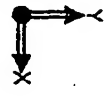
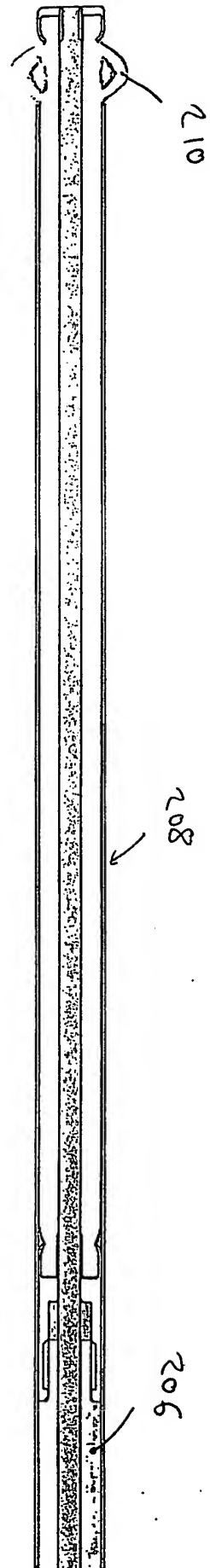


Fig 2B



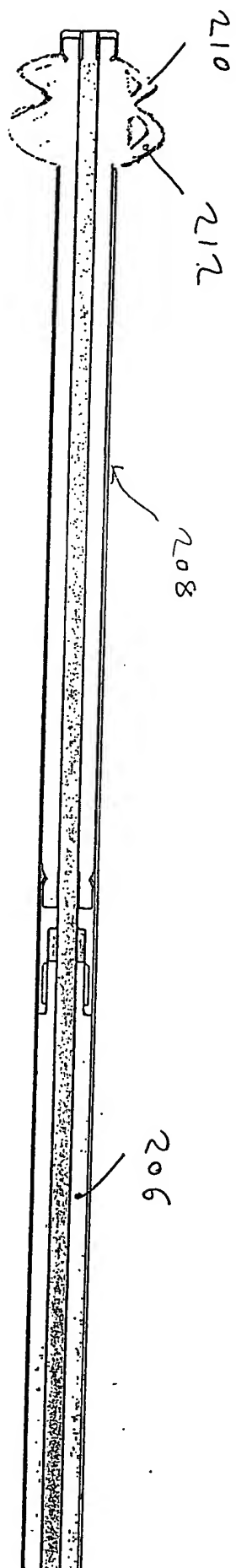


Fig 2C



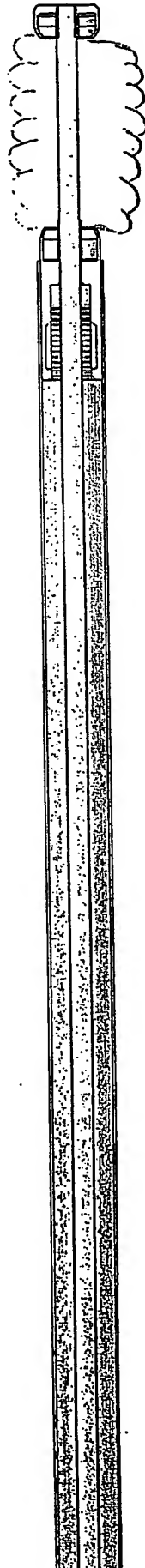


Fig.
2D

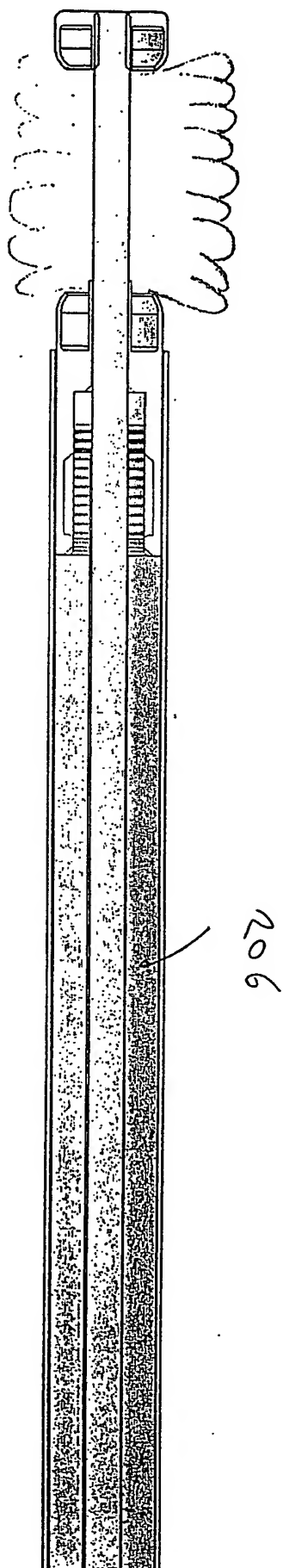


Fig. 2E



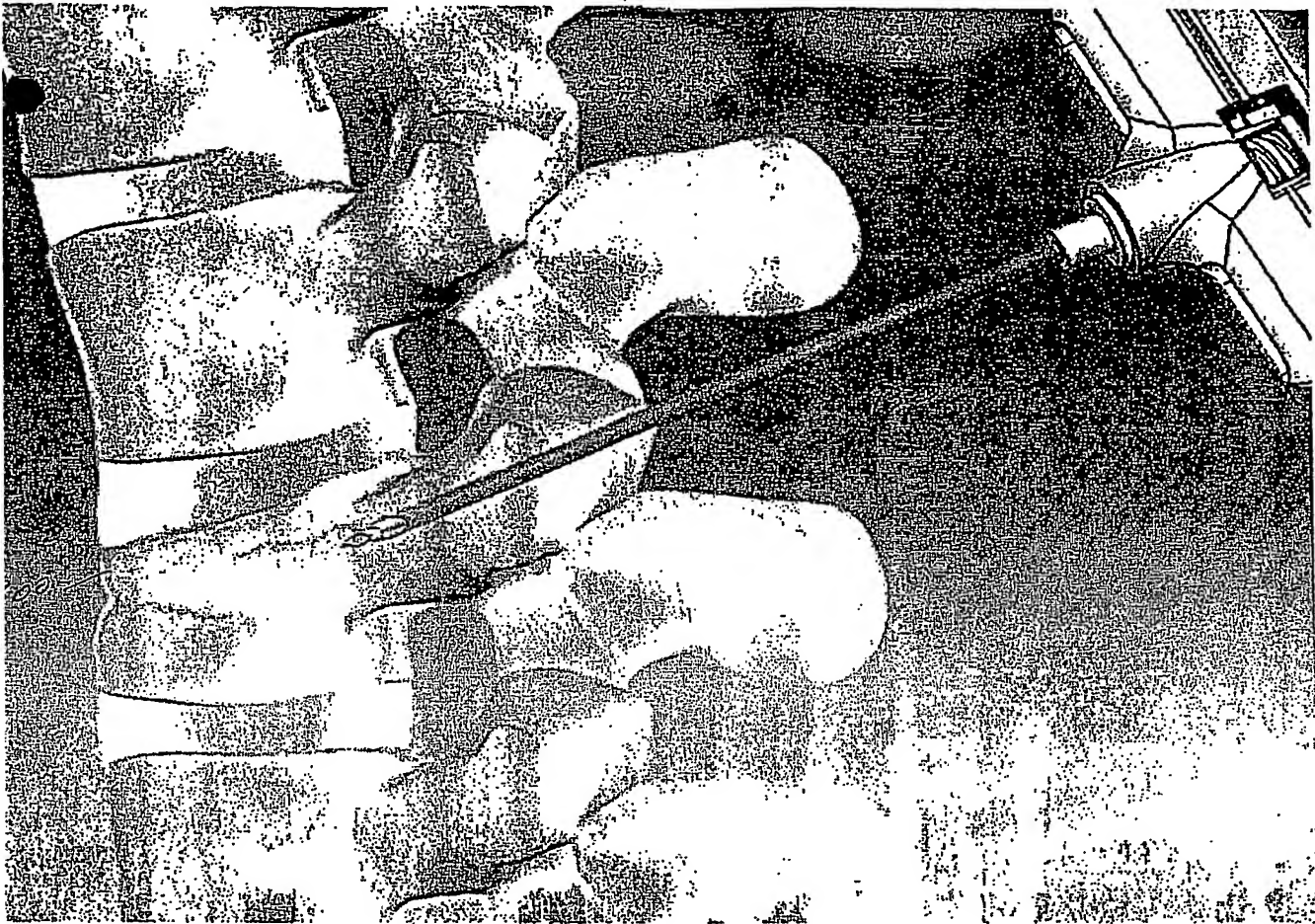


Fig 3A

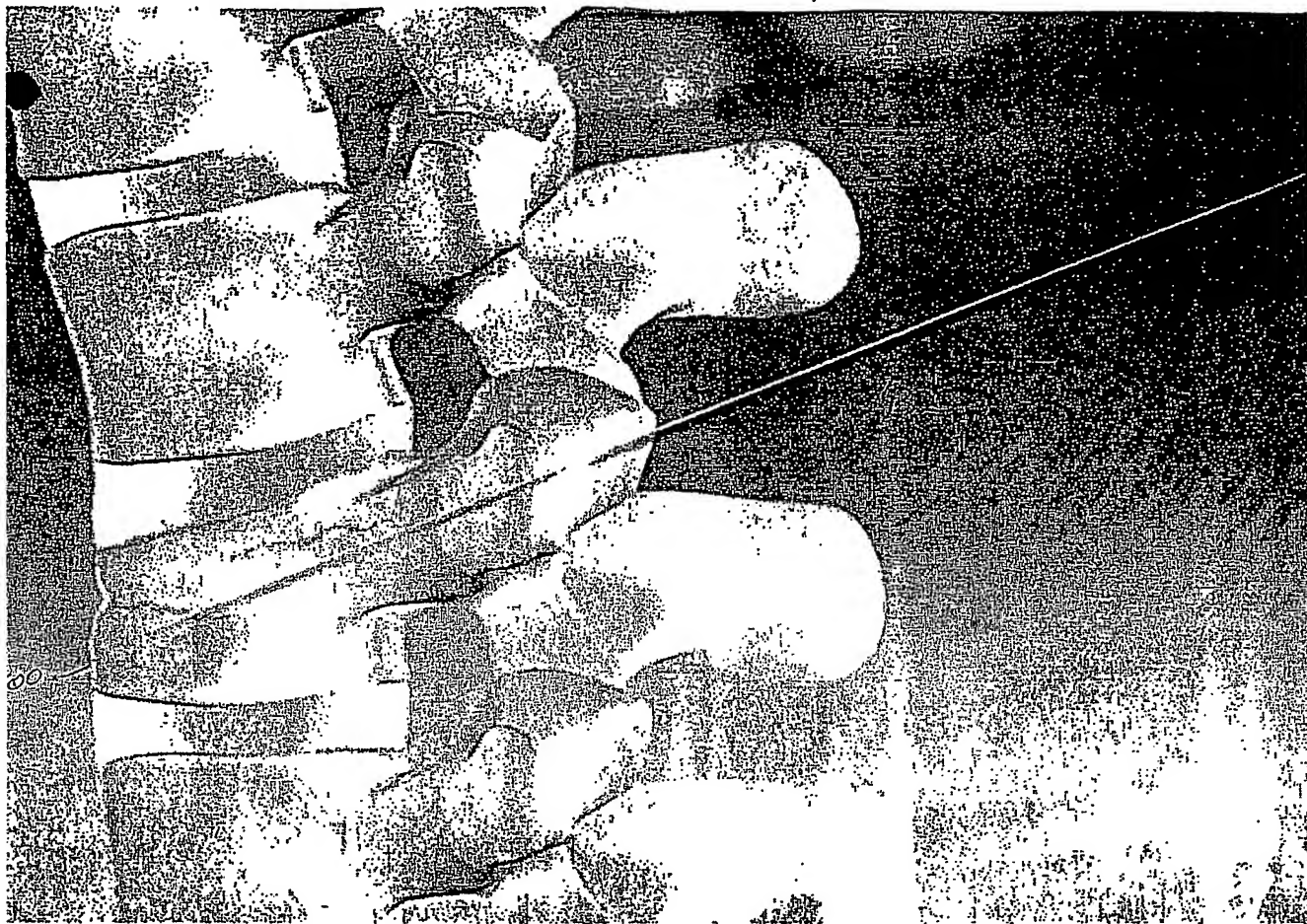


Fig 3B

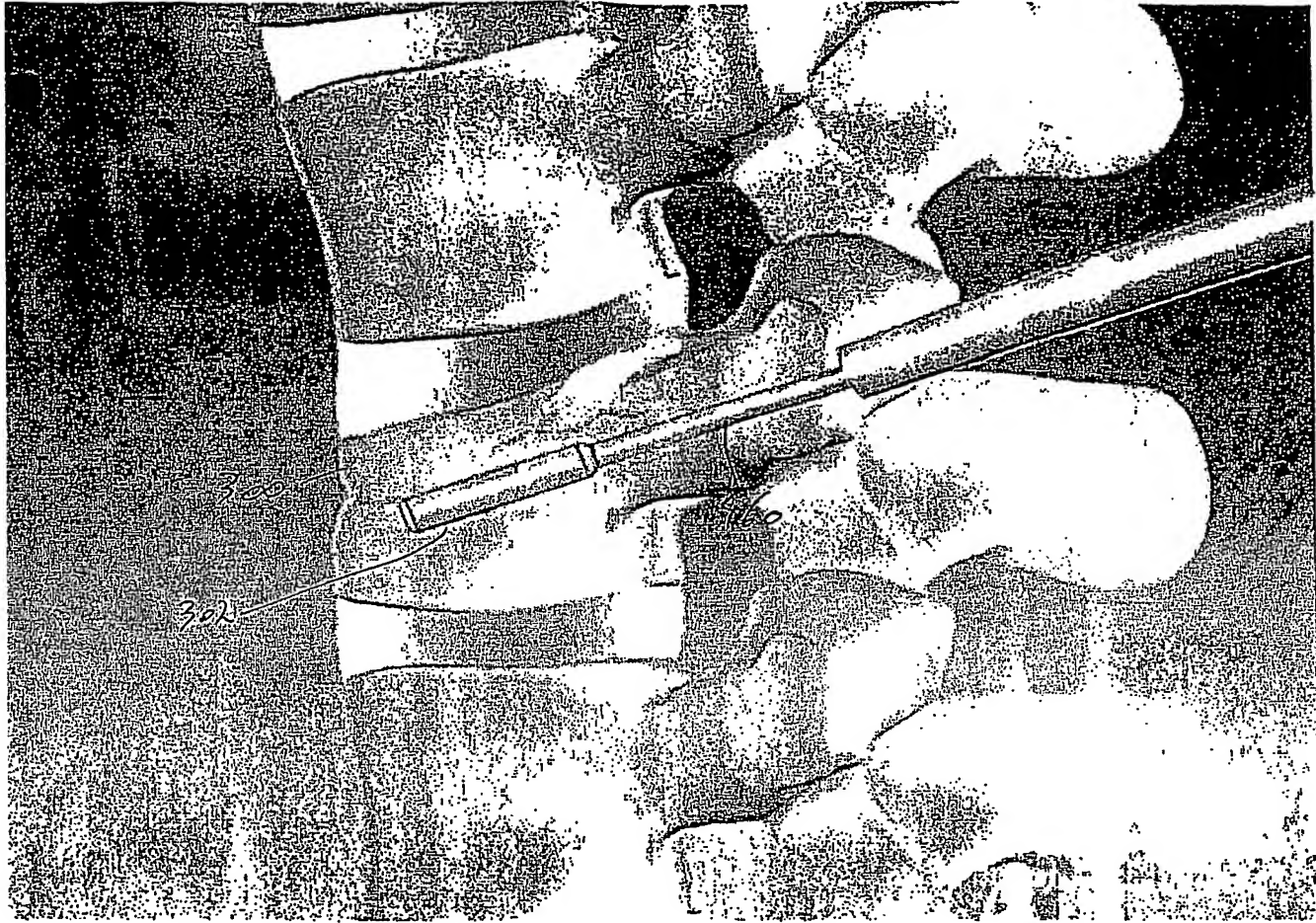


Fig. 3c

300
301

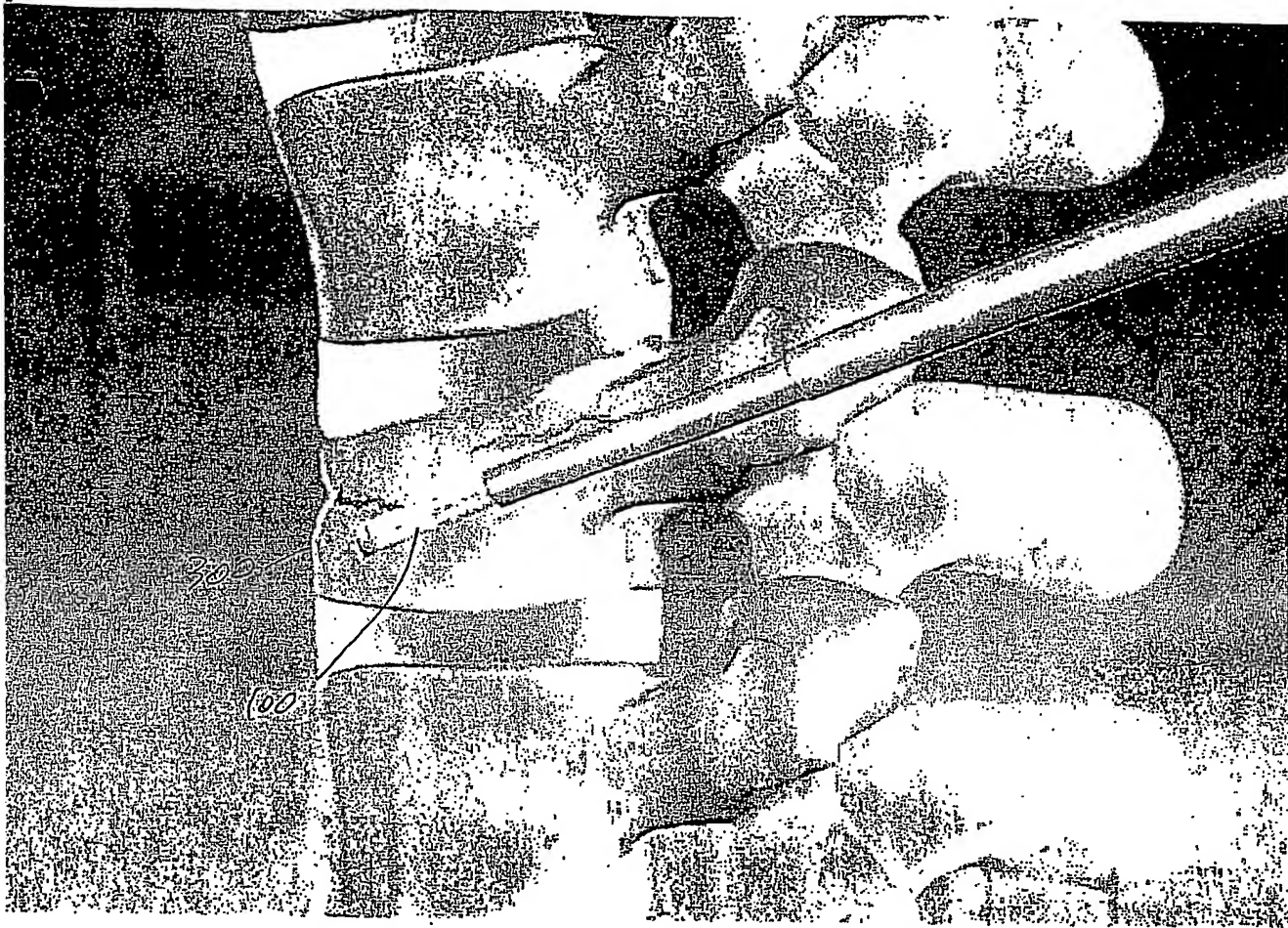


Fig. 3D
2.00

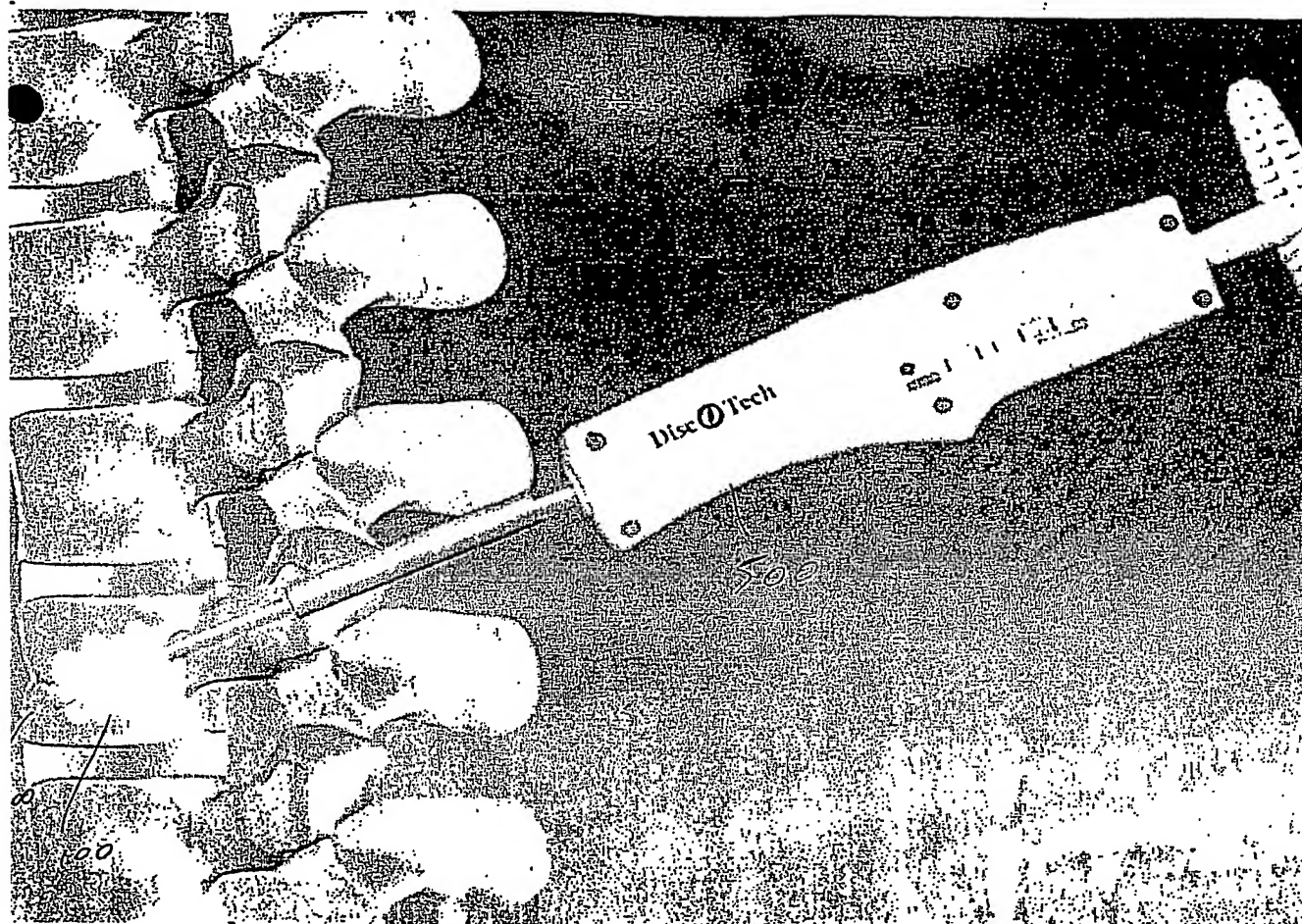


Fig 3E

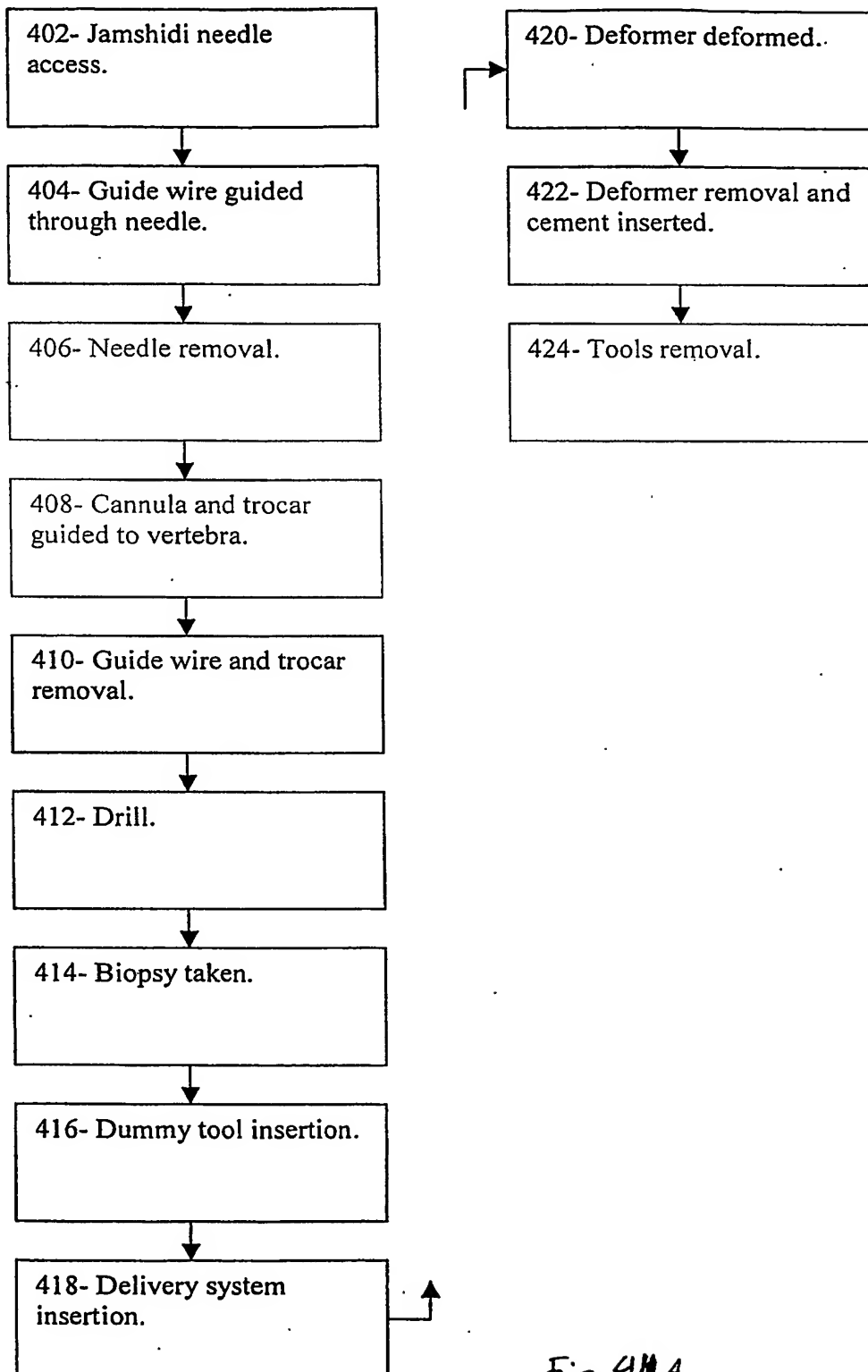


Fig. 4A

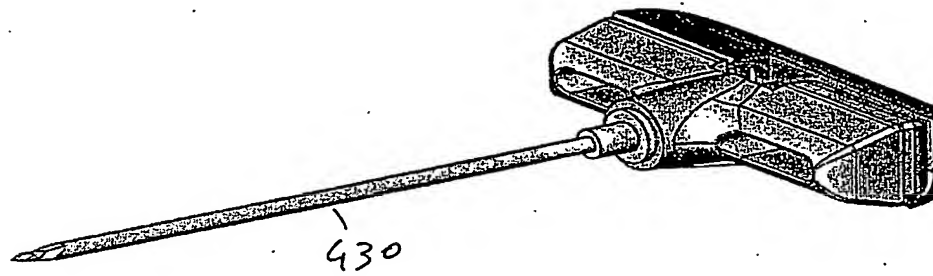


Fig. 4B

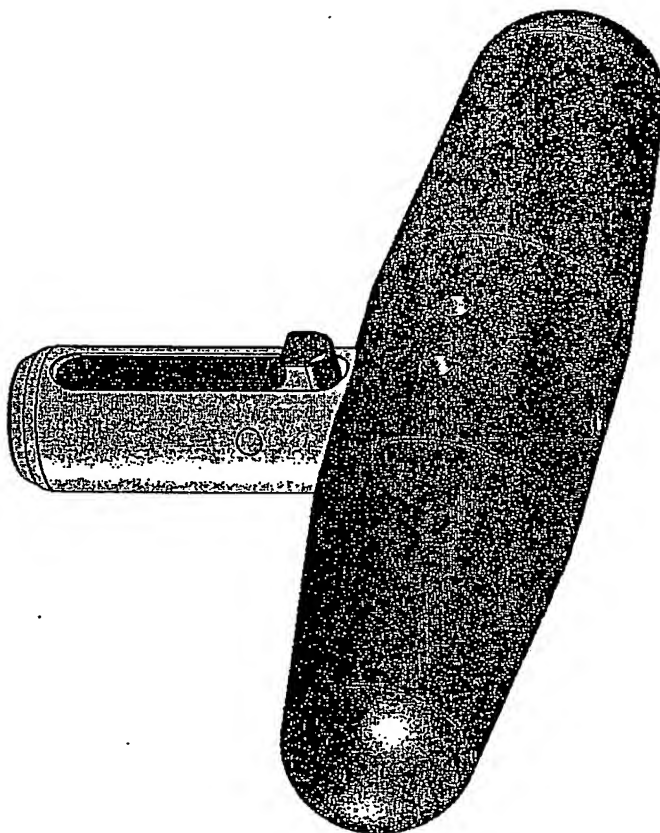


Fig. 4.C-1

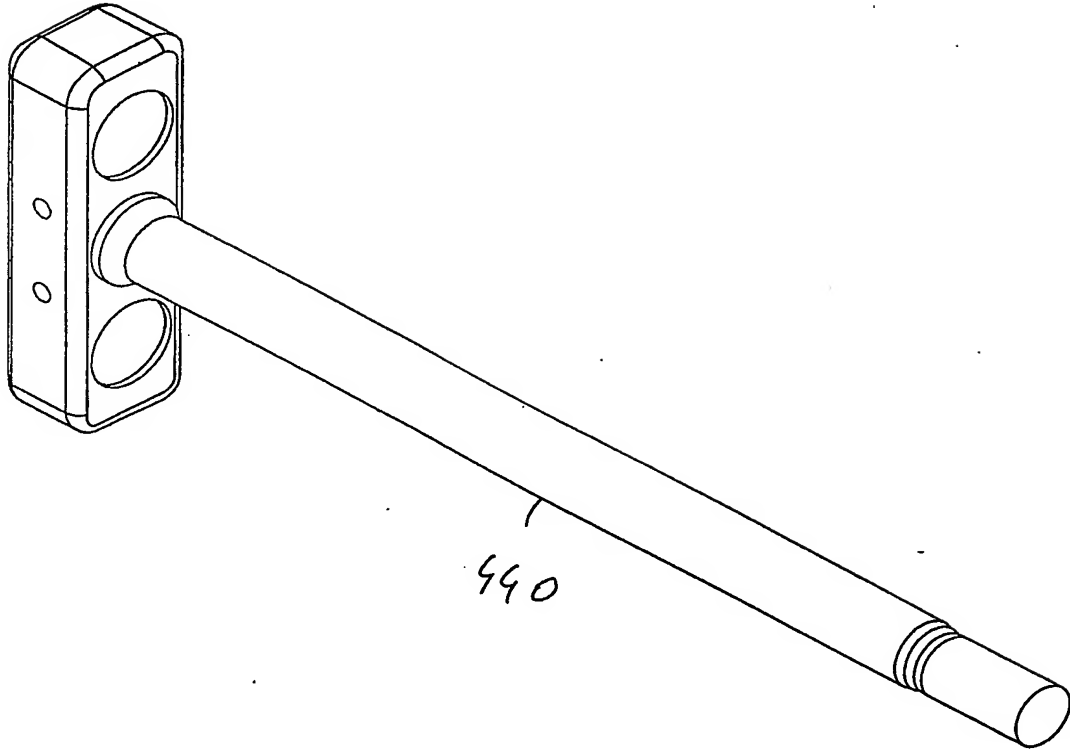


FIG. 4C

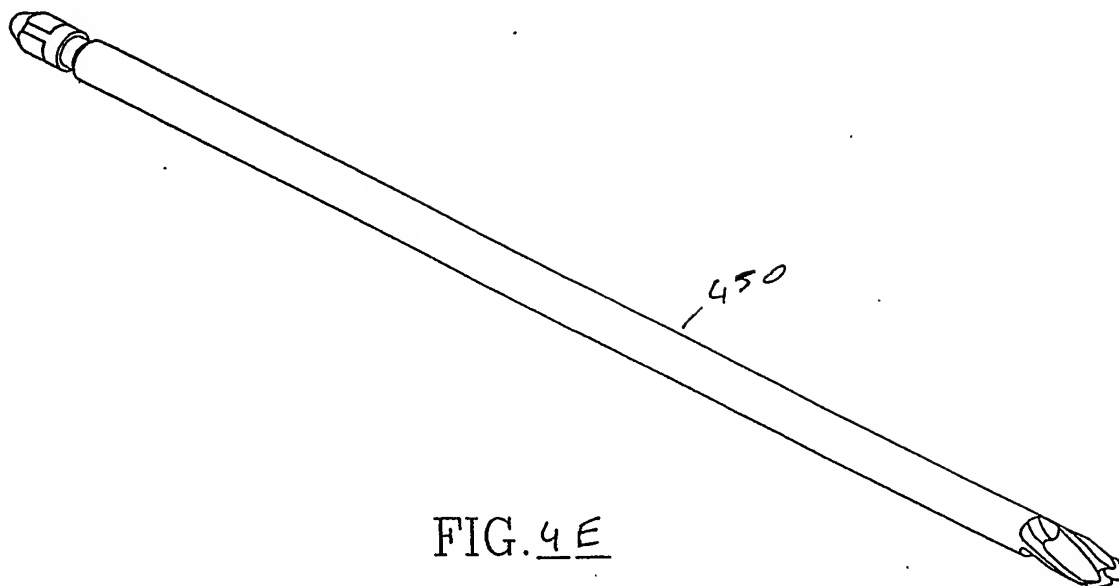


FIG. 4E

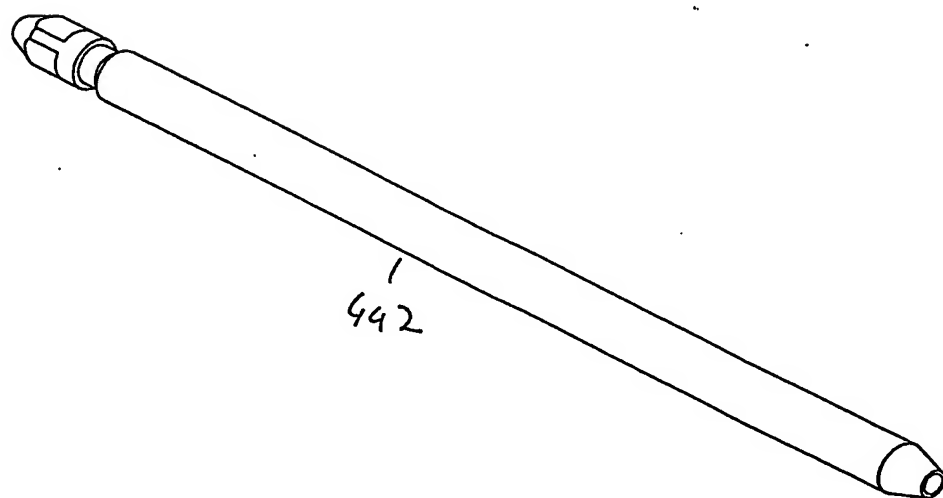


FIG. 4D

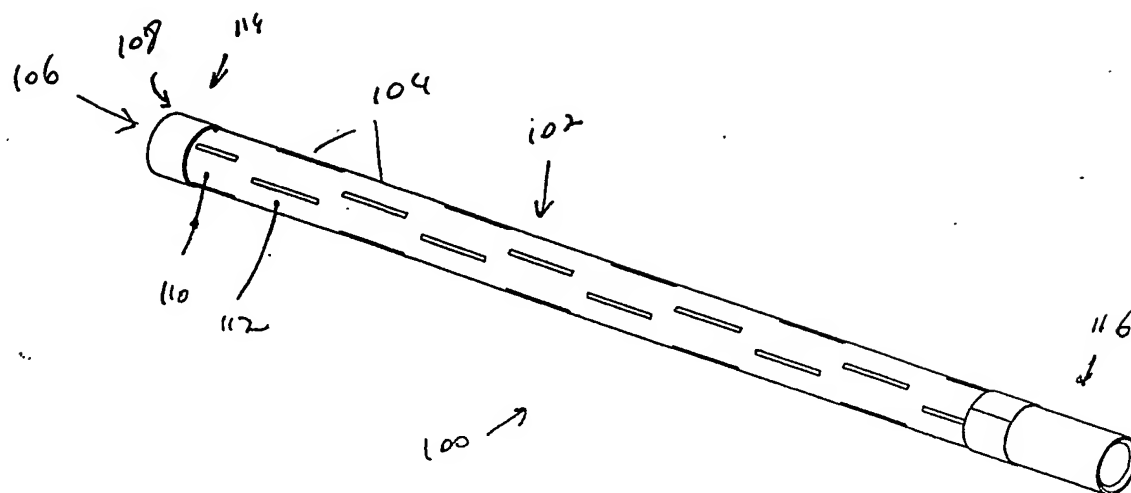


FIG. 1A

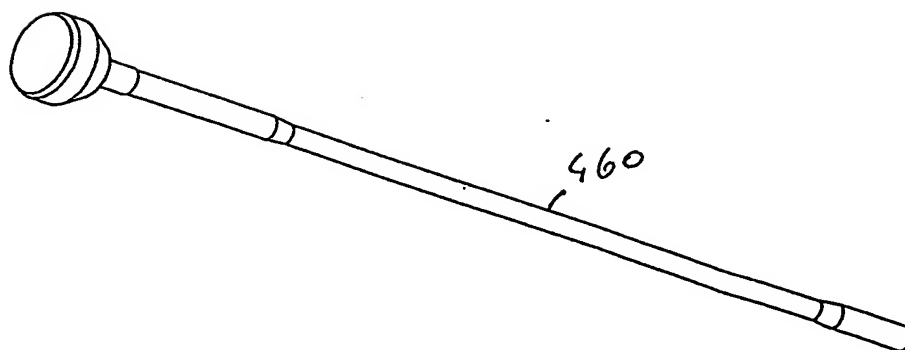


FIG. 4F

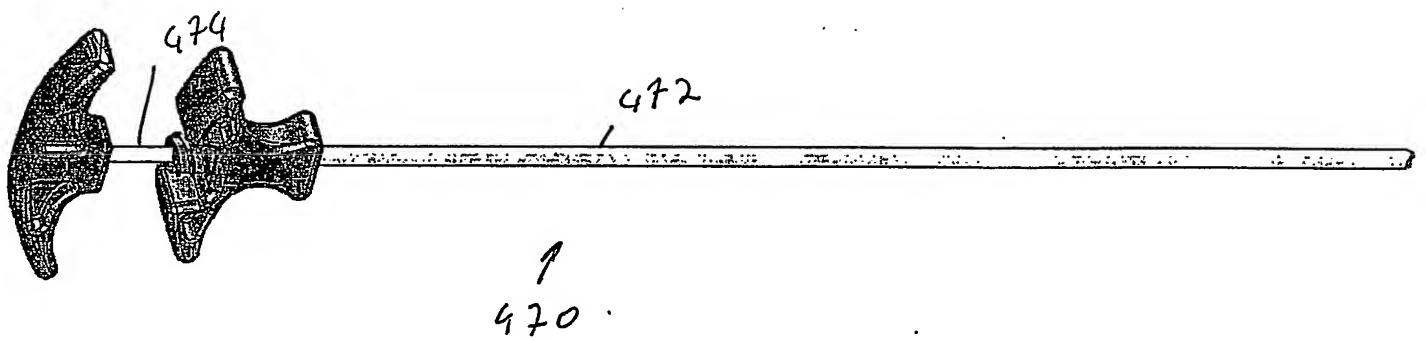
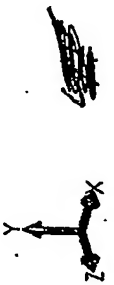
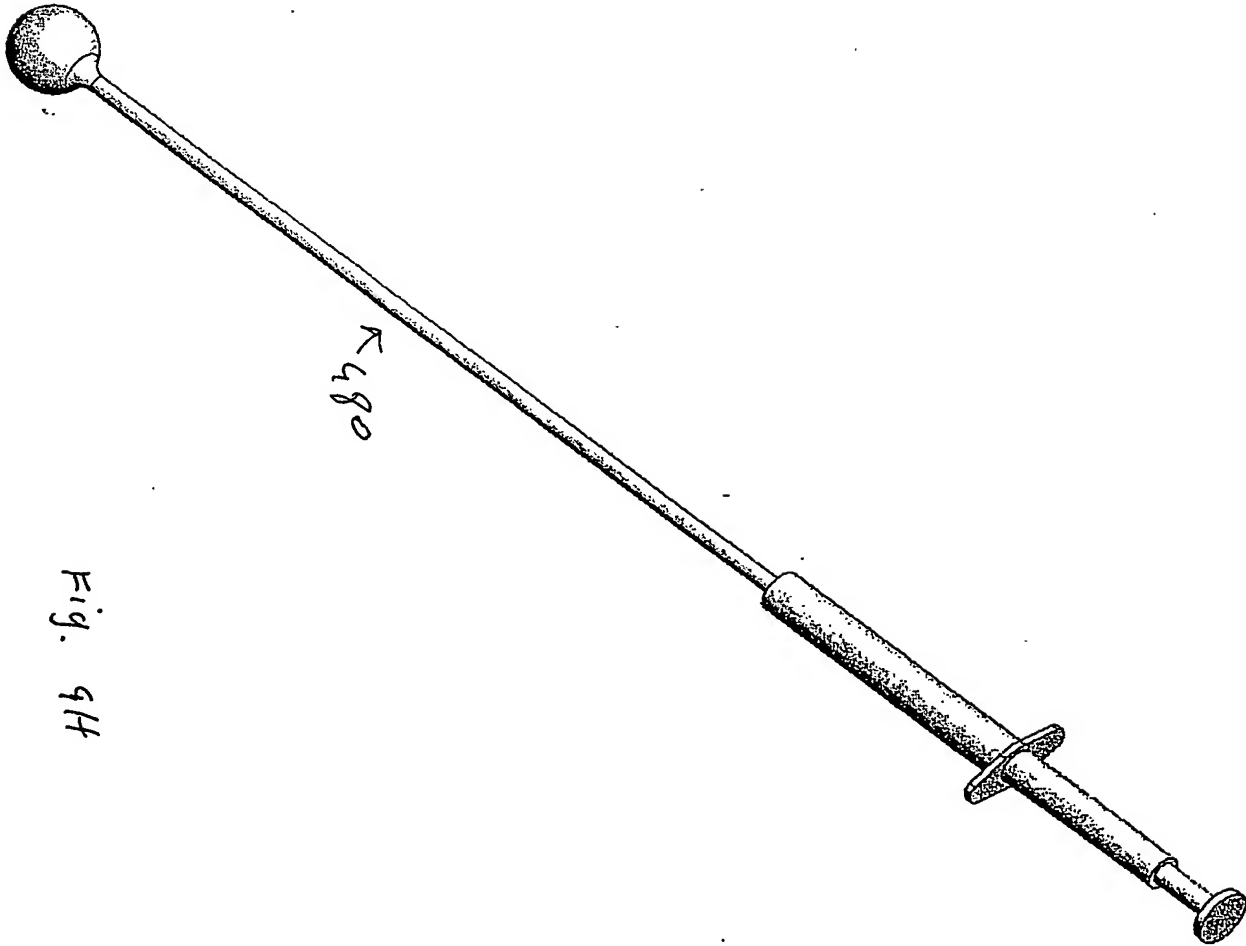


Fig. 4G





480-9H

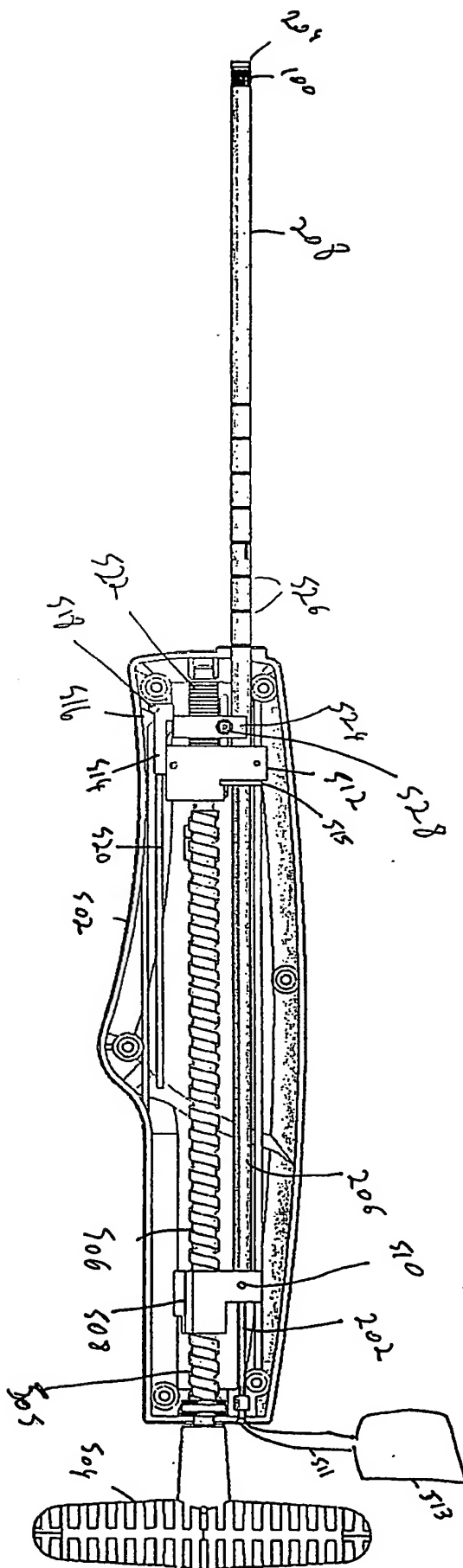
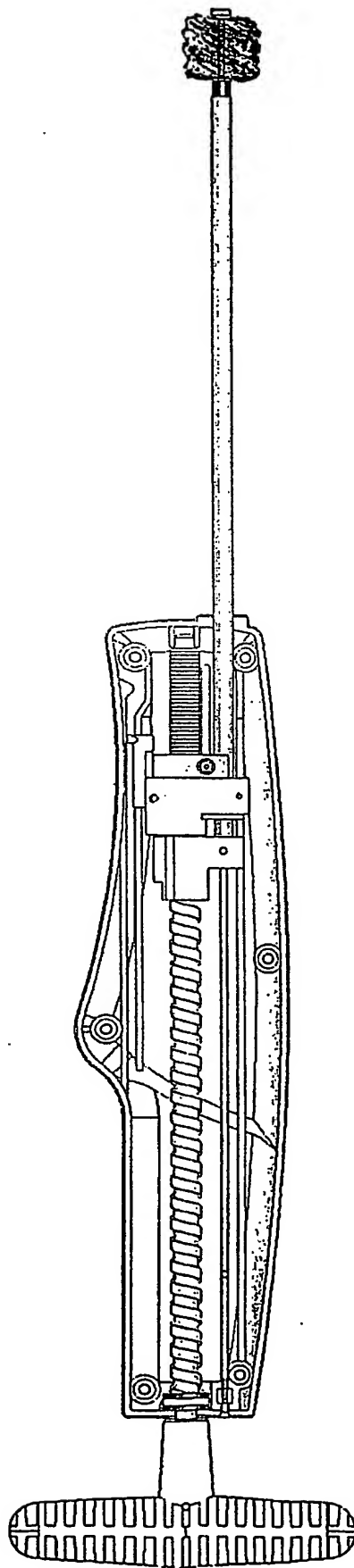


Fig. 5A

400



Fig 58



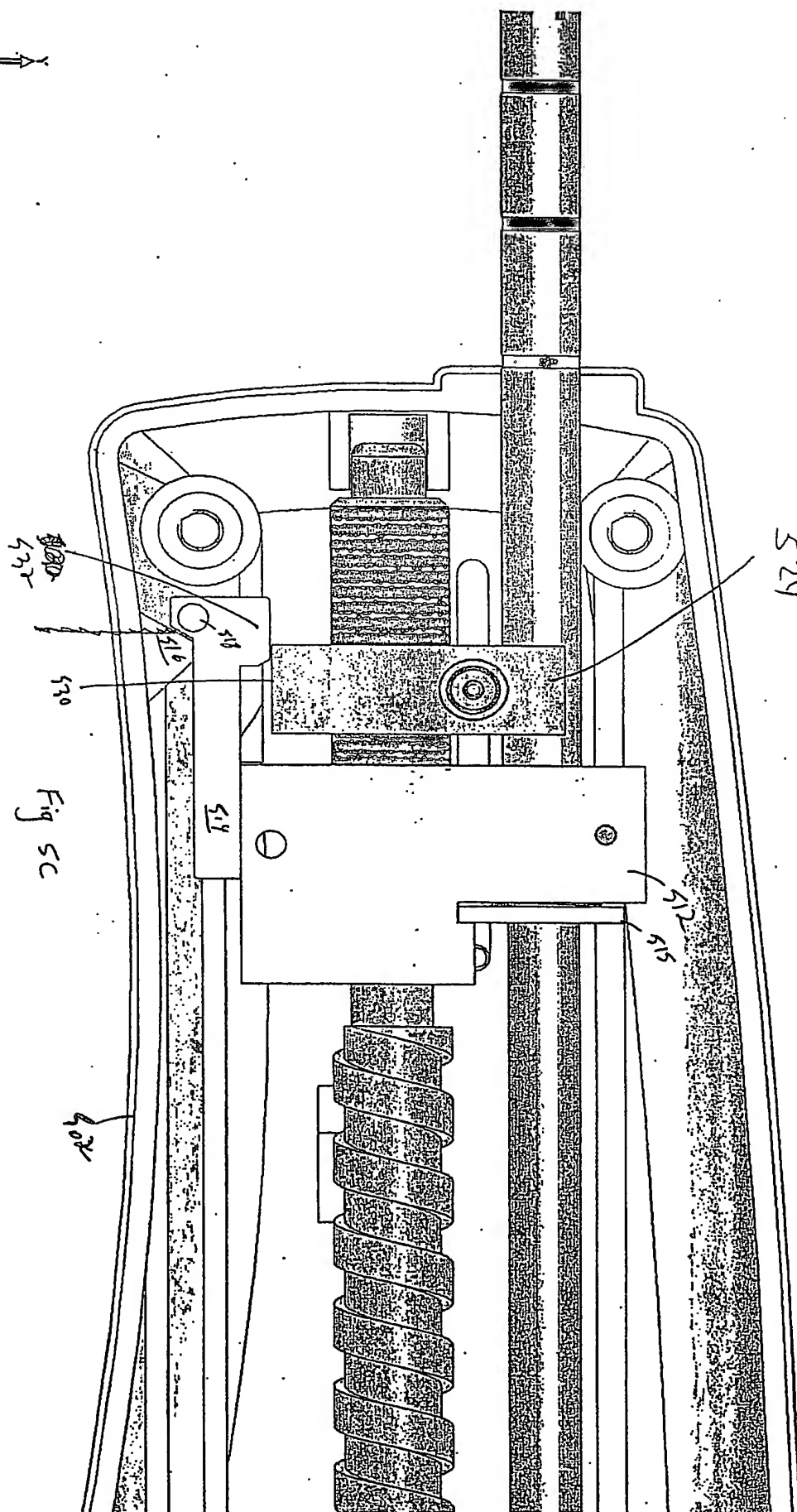
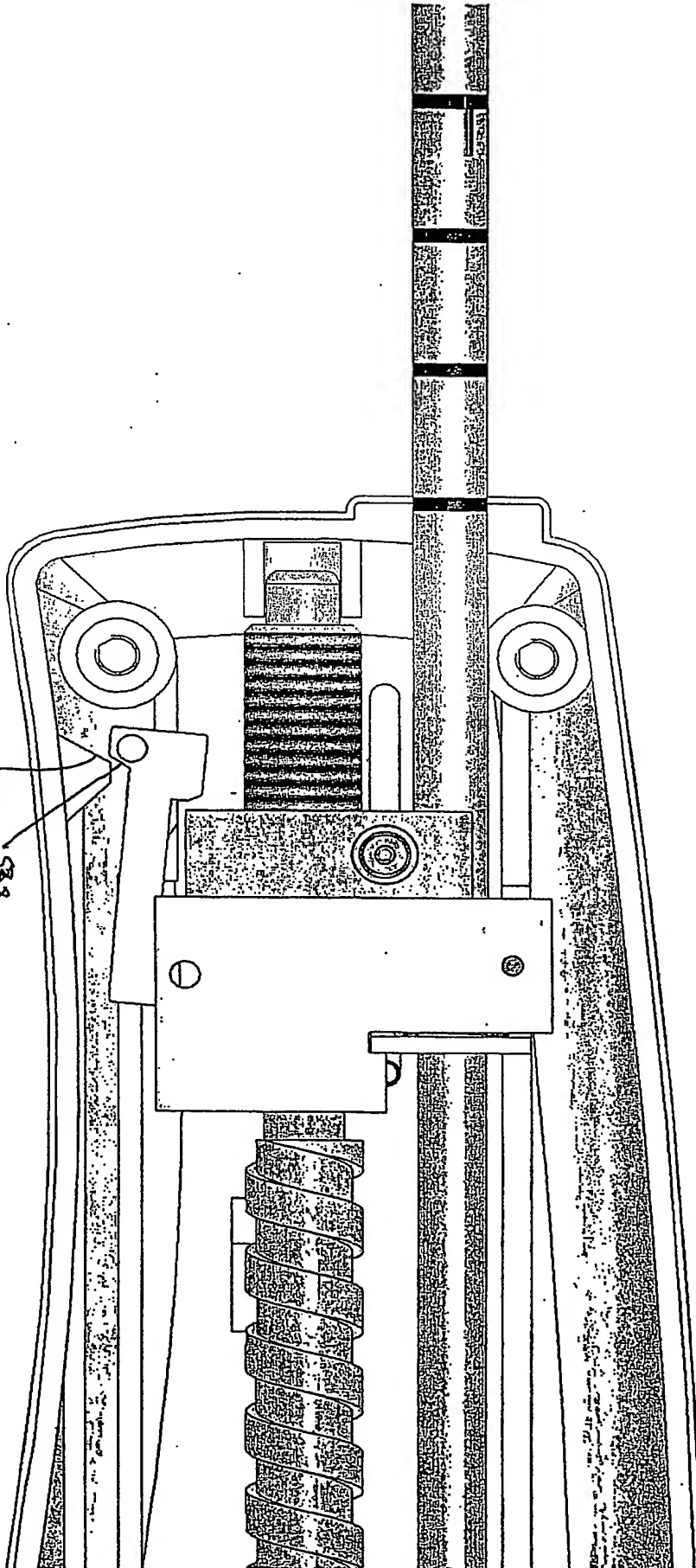




Fig 5D

536
538



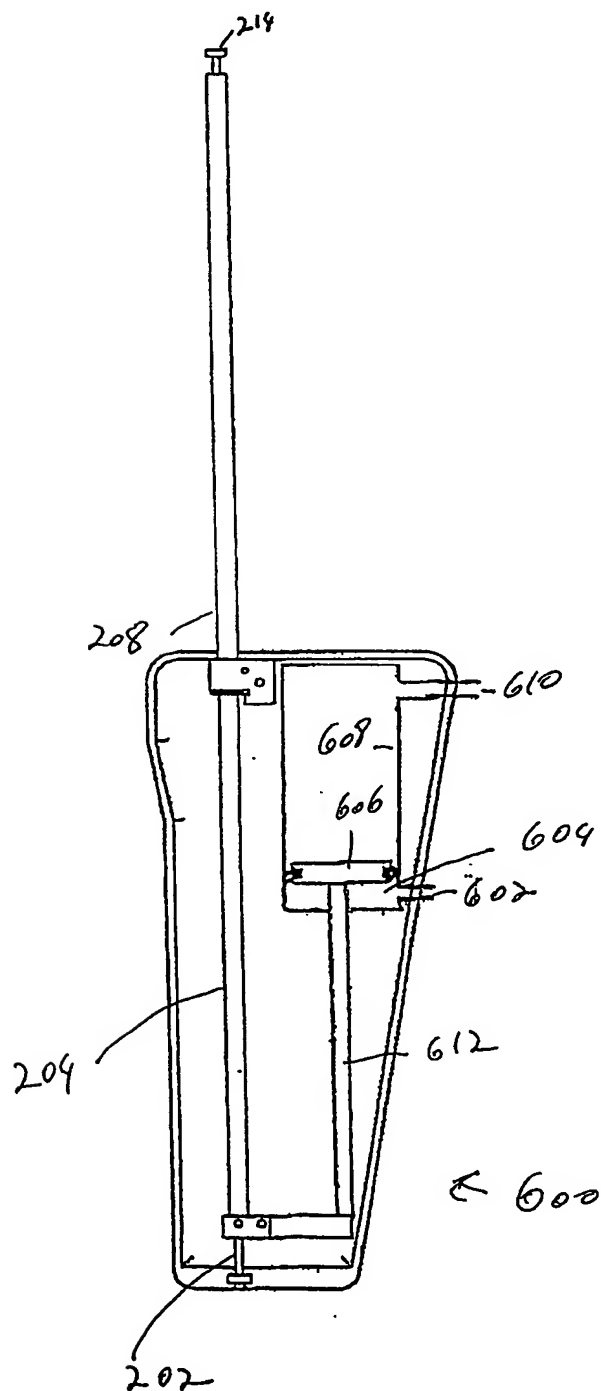
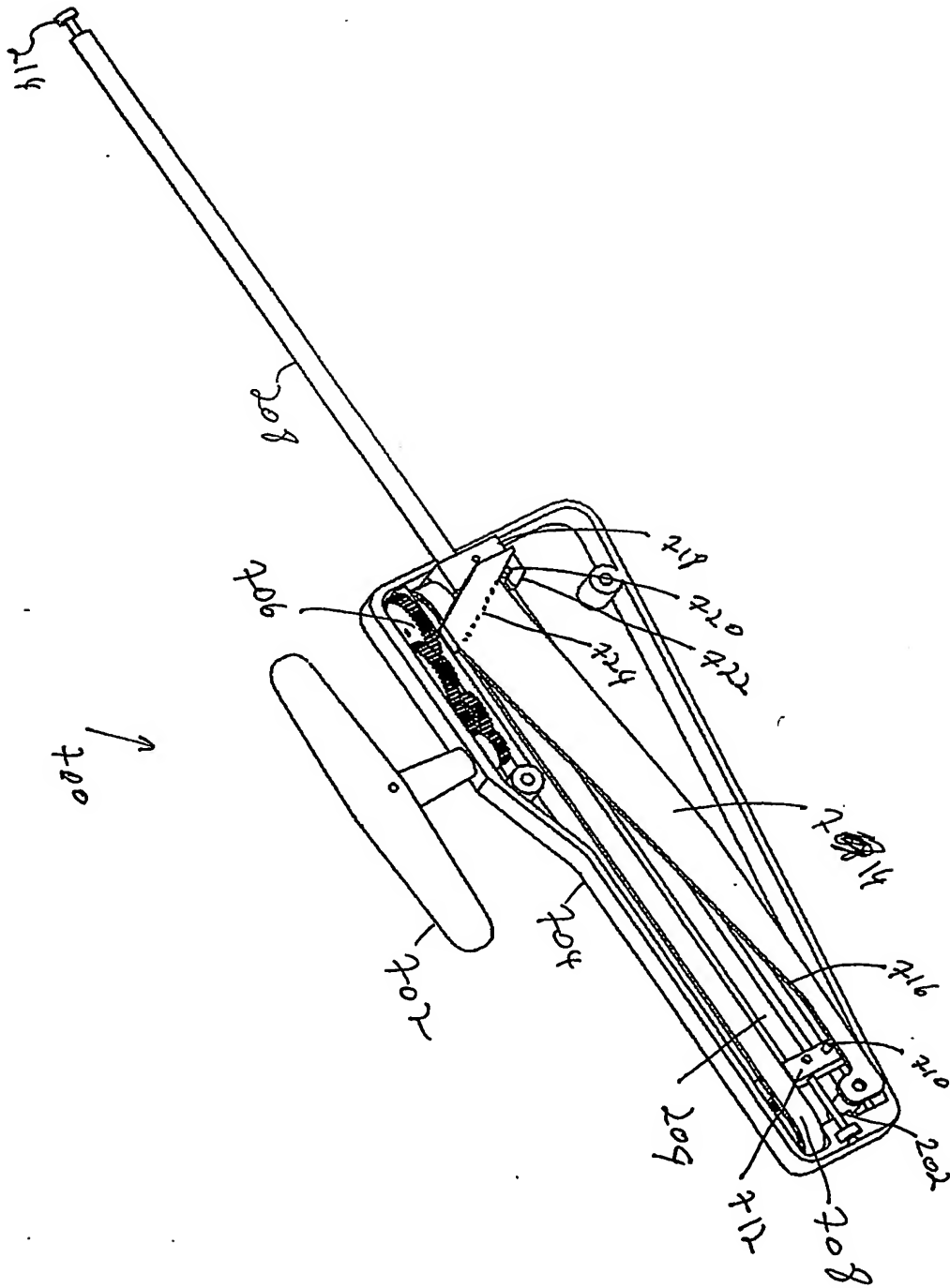


Fig 6



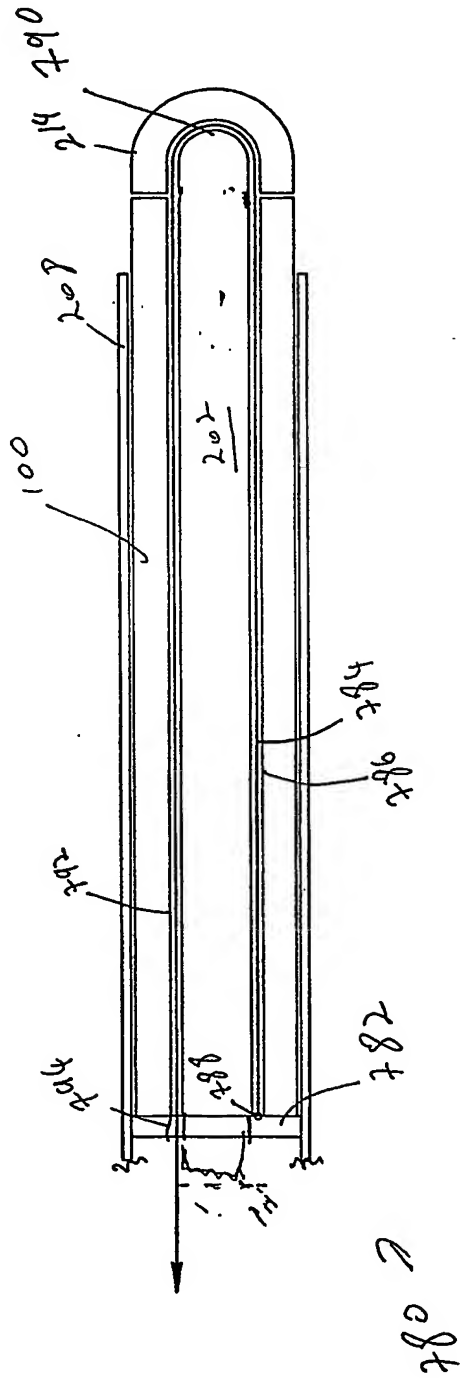


FIG. 7. B

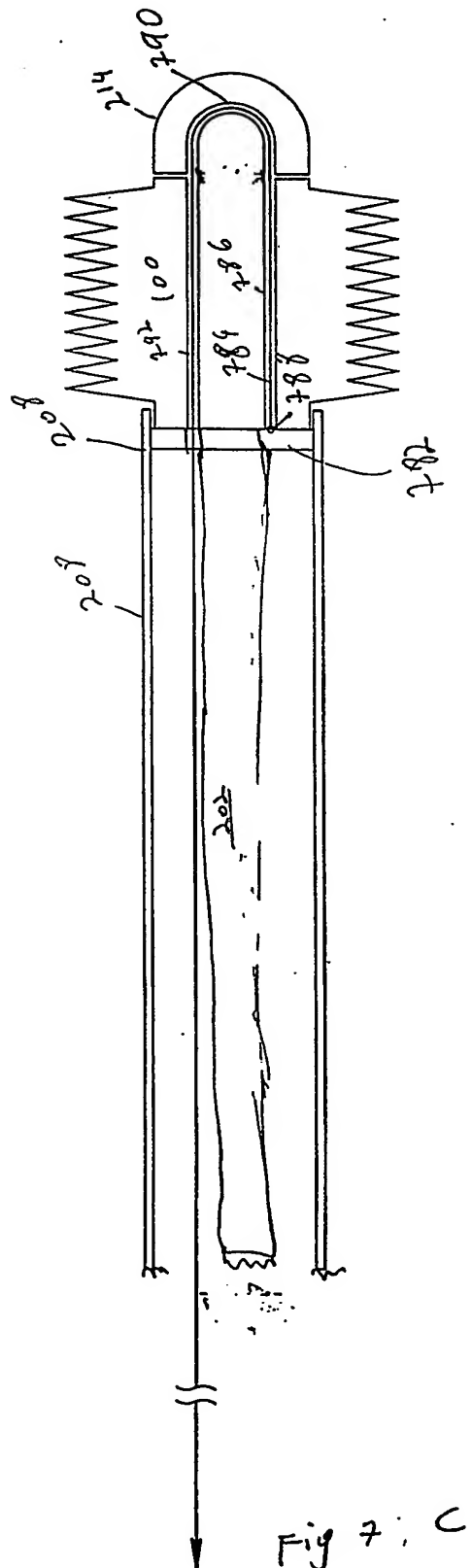


Fig 7, C

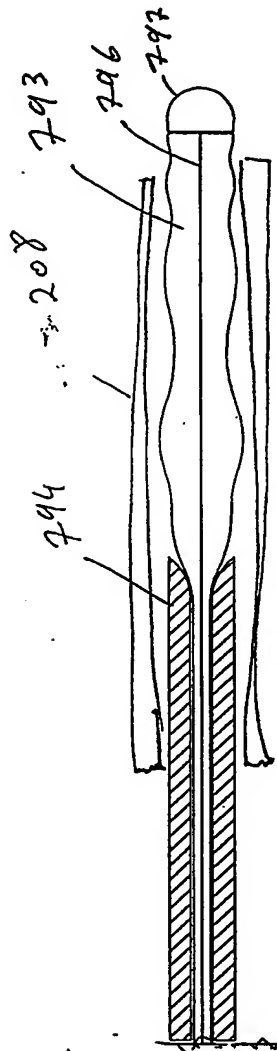


Fig 7D

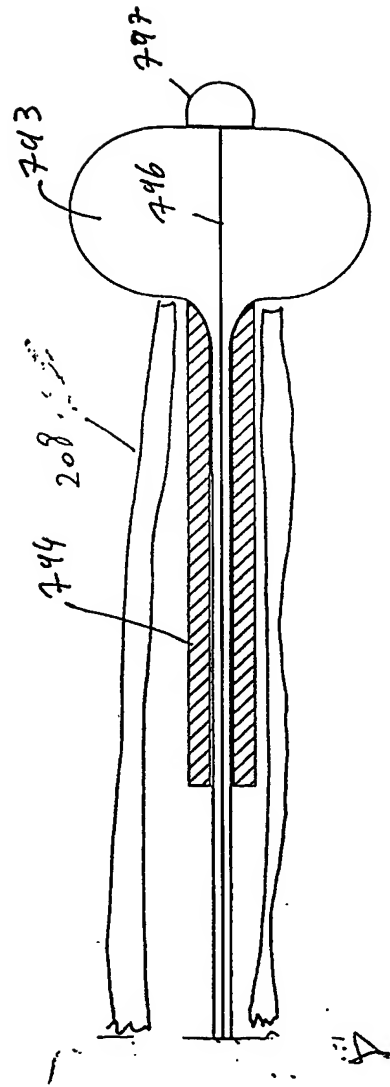


FIG. 7E

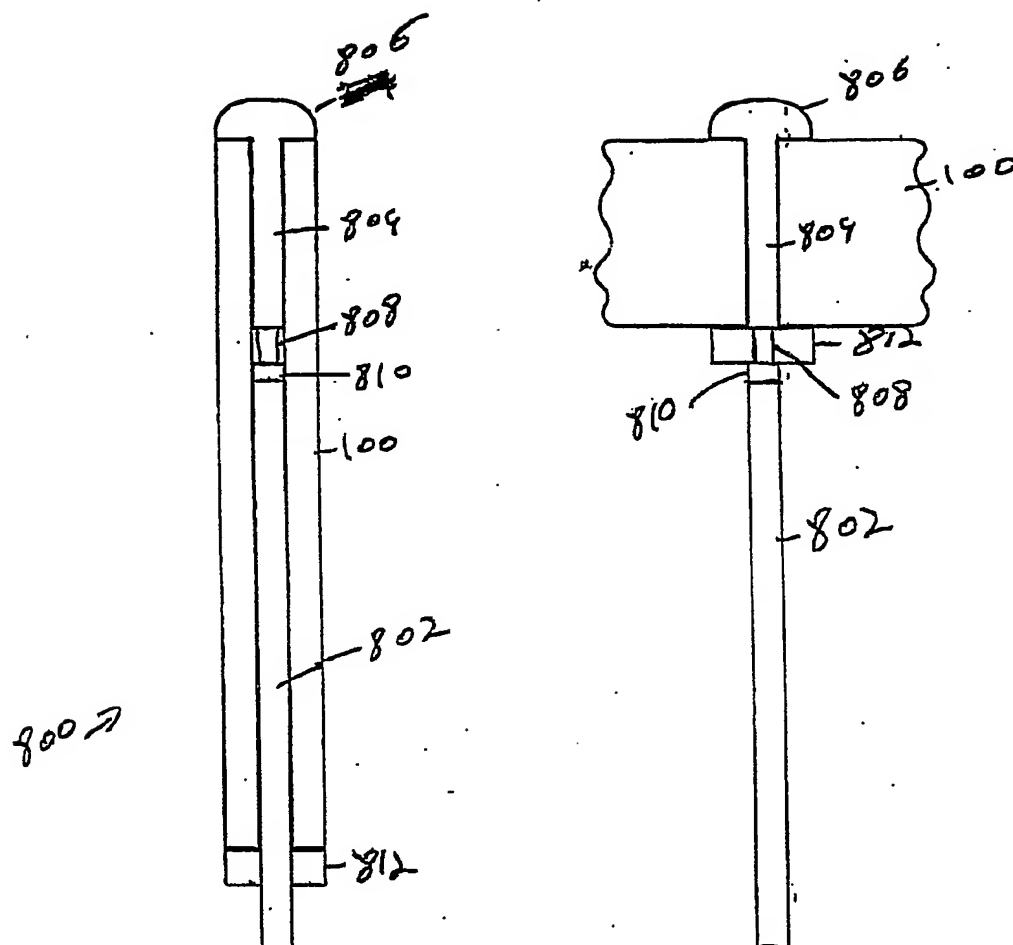


Fig 8A

~~FIG. 8(B)~~

Fig 8B

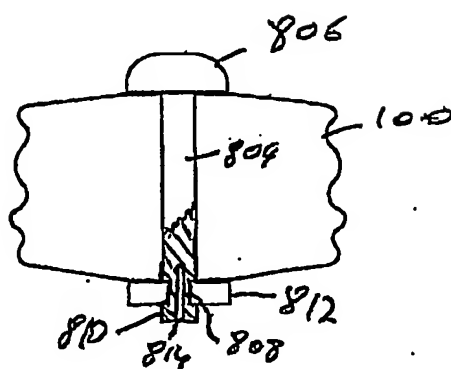


FIG. 8(C)

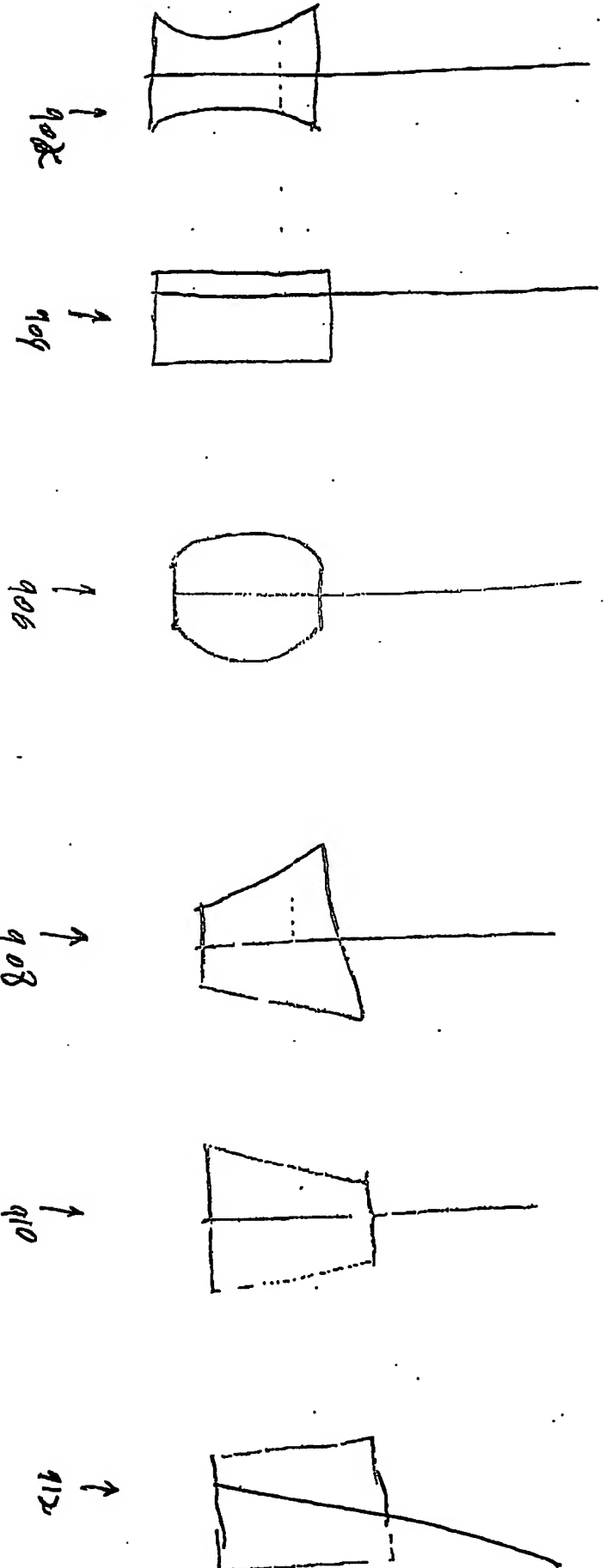
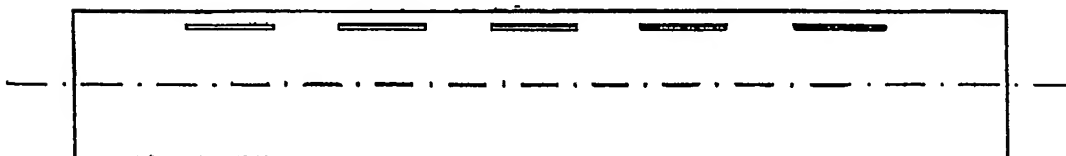
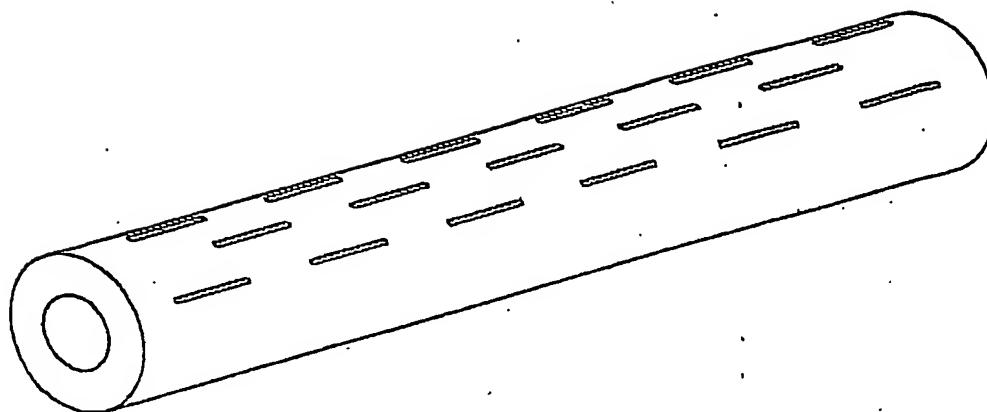


Fig. 9A



9/4 ↗

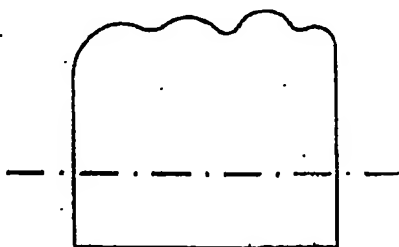


FIG.9b

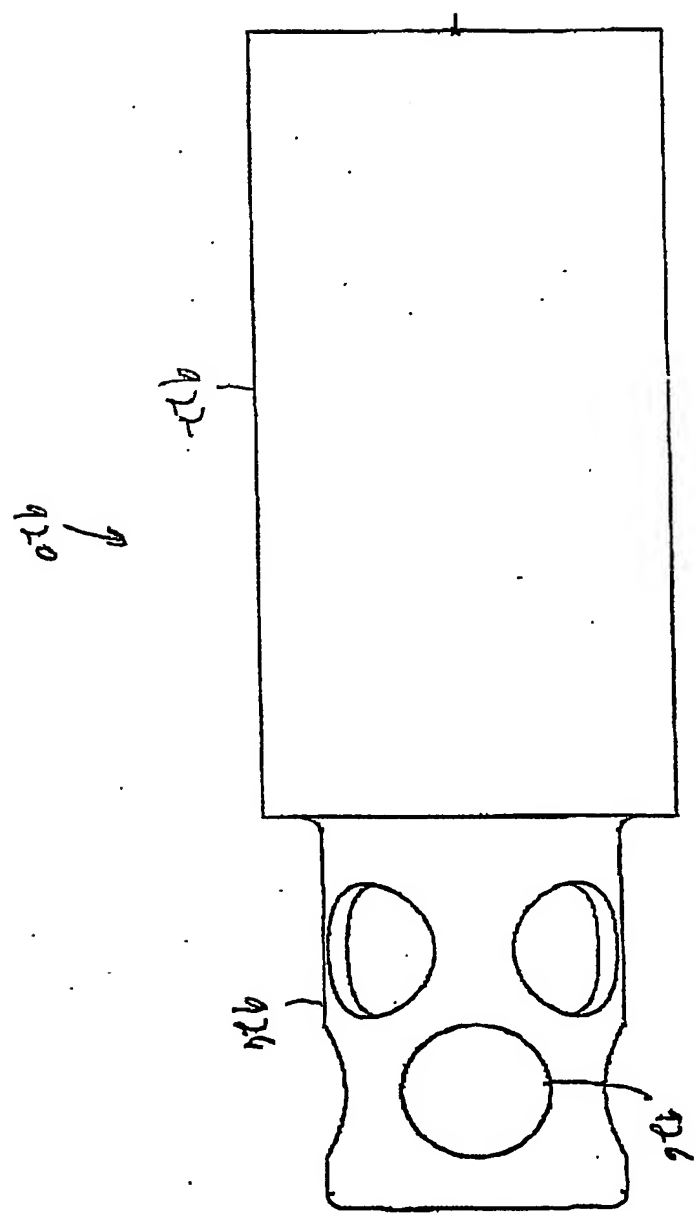
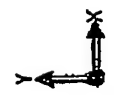


Fig. 9C



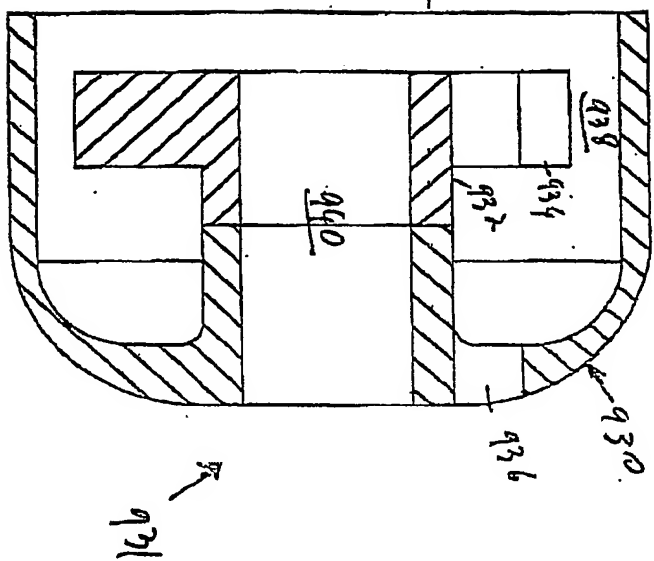


Fig. 9D

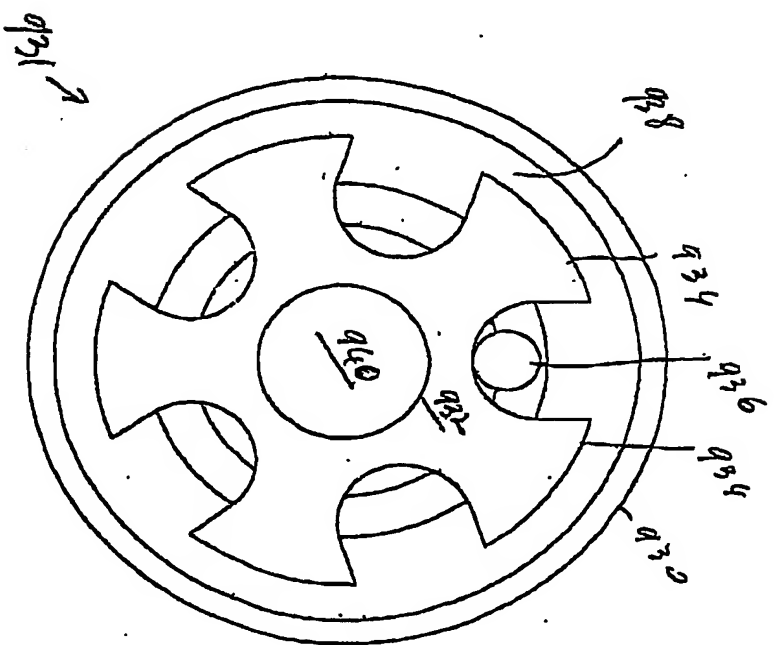


Fig. 9E

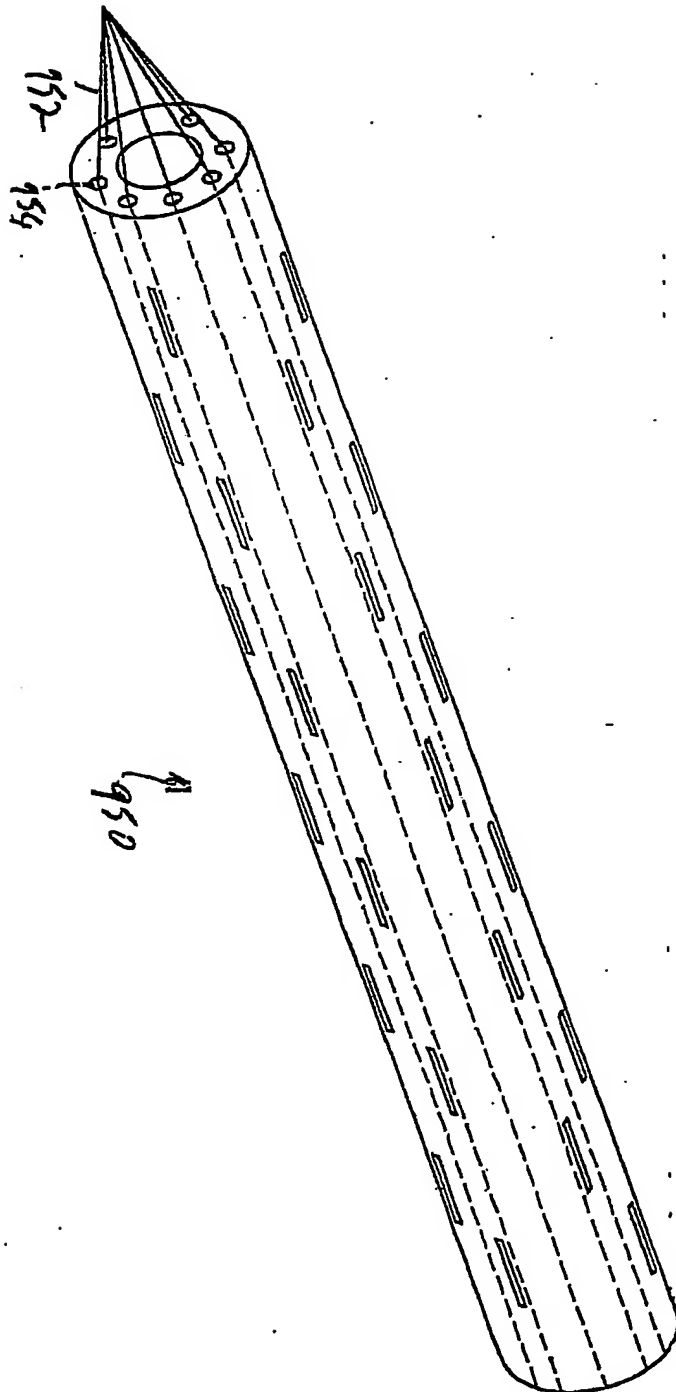


FIG. 9

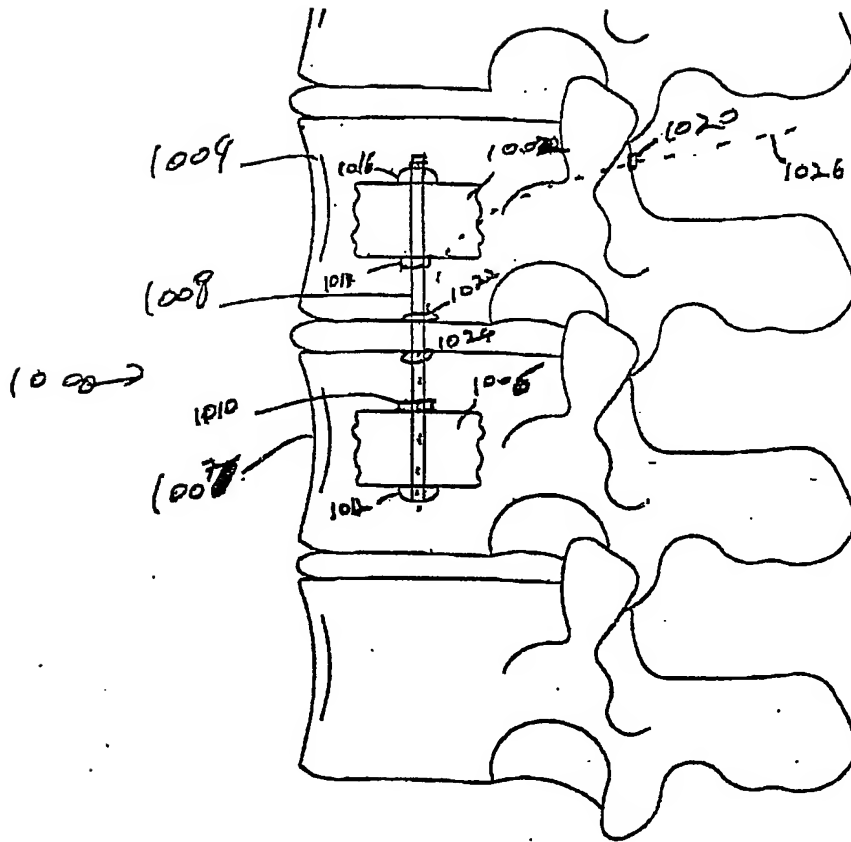


FIG.10

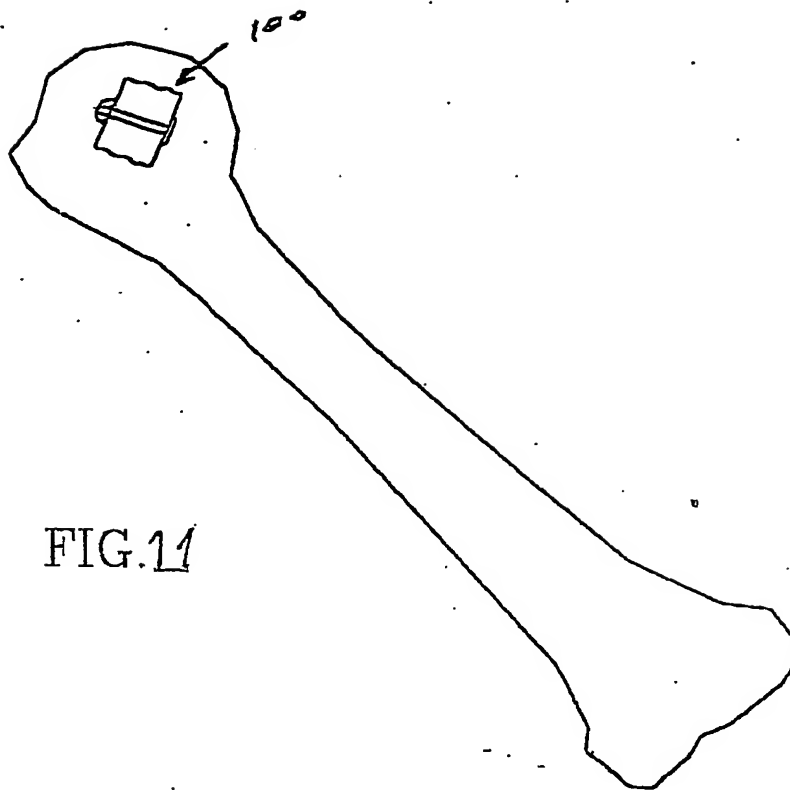


FIG. 11

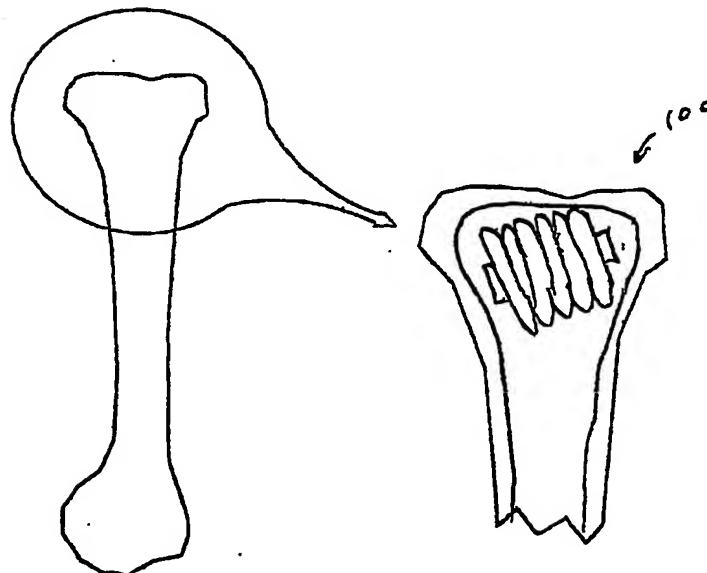
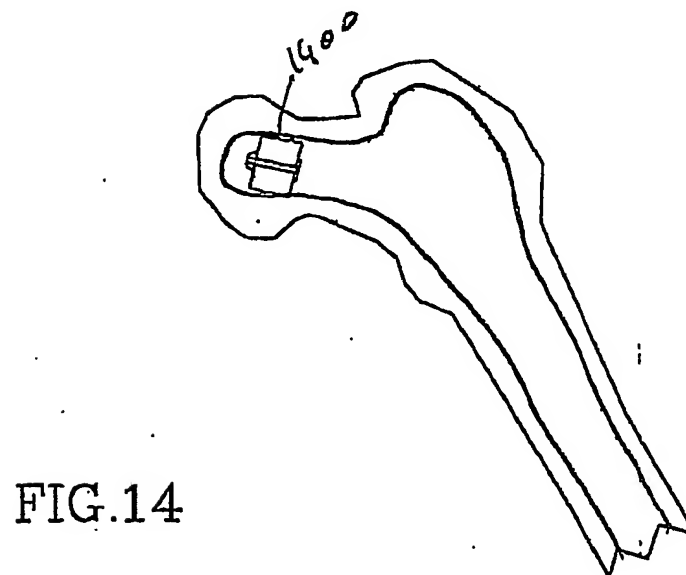
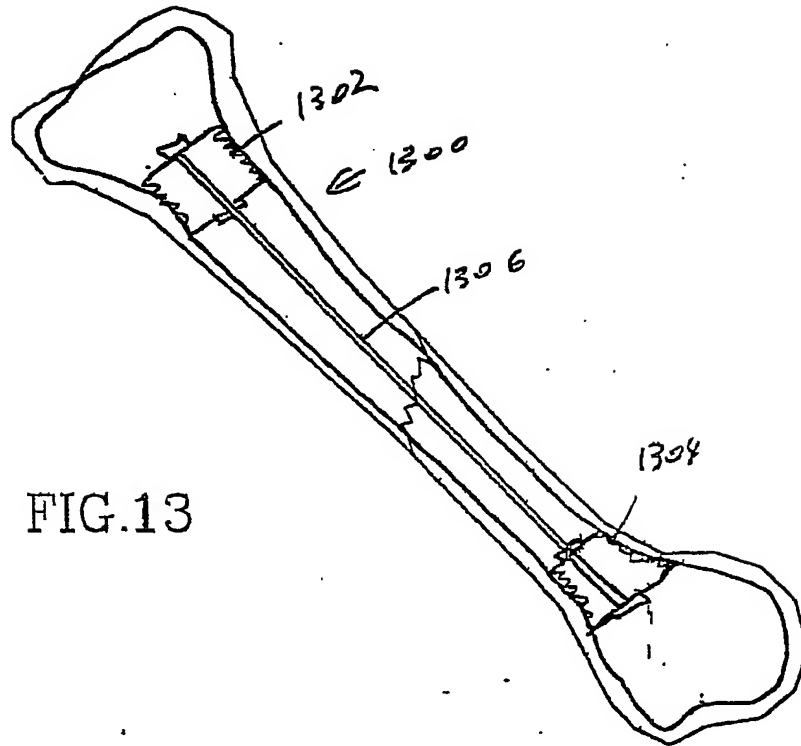
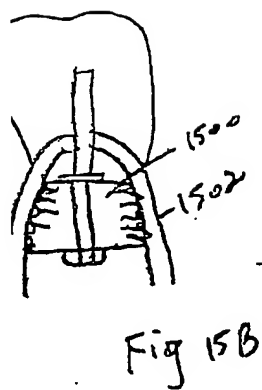
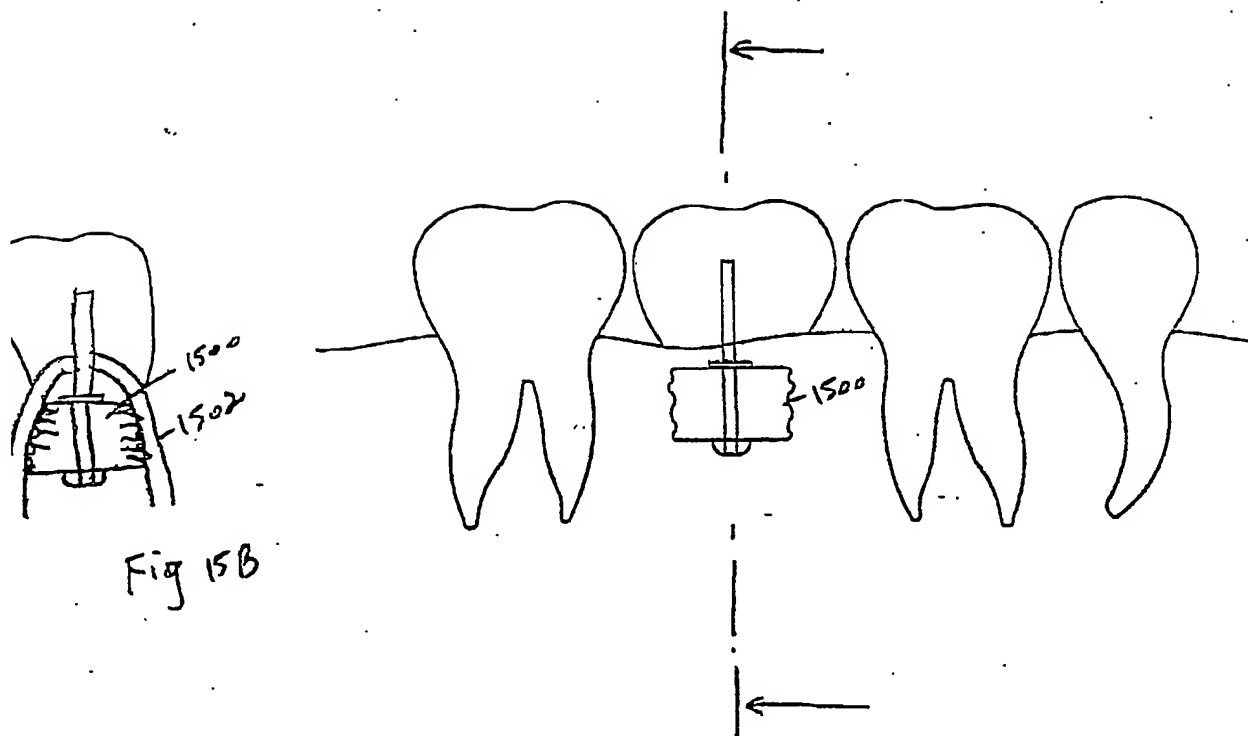


FIG. 12





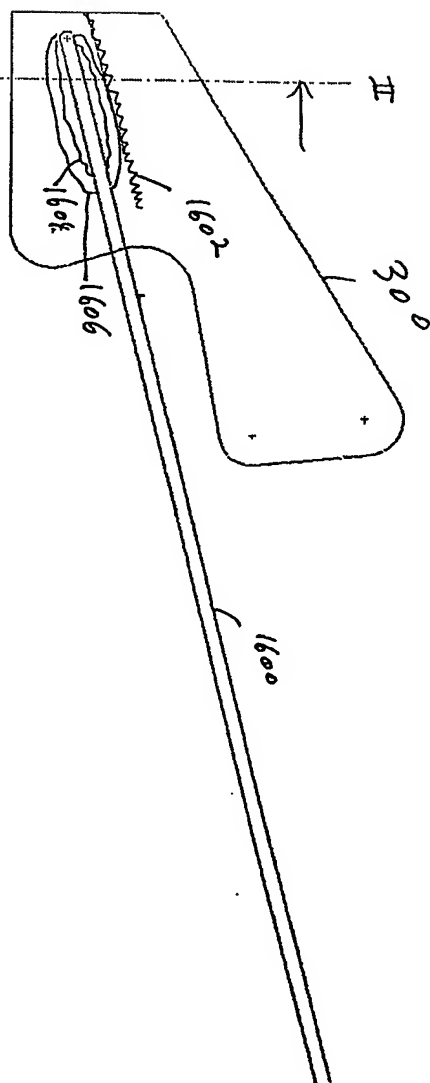


Fig. 16A

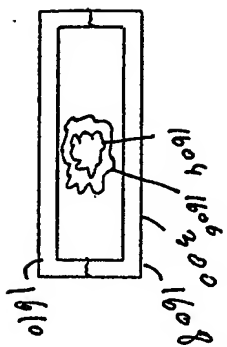


Fig. 16B

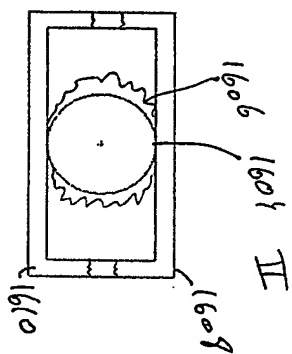


Fig. 16C

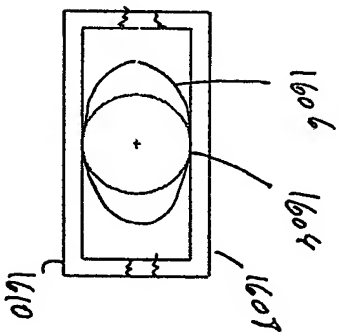


Fig. 16D

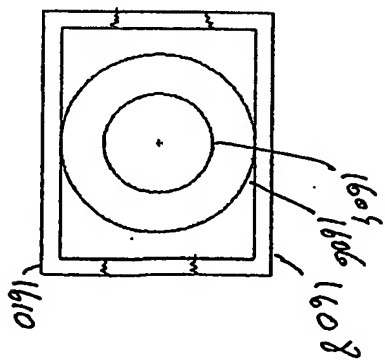


Fig. 16E

**This Page is Inserted by IFW Indexing and Scanning
Operations and is not part of the Official Record**

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

- ☐ **BLACK BORDERS**
- ☐ **IMAGE CUT OFF AT TOP, BOTTOM OR SIDES**
- ☐ **FADED TEXT OR DRAWING**
- ☐ **BLURRED OR ILLEGIBLE TEXT OR DRAWING**
- ☐ **SKEWED/SLANTED IMAGES**
- ☐ **COLOR OR BLACK AND WHITE PHOTOGRAPHS**
- ☐ **GRAY SCALE DOCUMENTS**
- ☐ **LINES OR MARKS ON ORIGINAL DOCUMENT**
- ☐ **REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY**
- ☐ **OTHER:** _____

IMAGES ARE BEST AVAILABLE COPY.

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.